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The Workshop on ML and its Applications was held
June 20-21, 1992 in San Francisco, CA, in conjunction with the
ACM SIGPLAN '92 Conferences and Workshops.

Abstract

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implementation. As the language and its implementations have matured, the range of
applications has increased greatly. New applications, combined with new ideas in design and
implementation, have stimulated a large number of significant activities in research and software
development. This workshop, like the previous workshops in Princeton, Edinburgh, and
Pittsburgh, provided a forum for theses activities, with a special emphasis on applications of the
language. A total of 21 abstracts were selected for presentation and discussion at the workshop:
12 were chosen for conference-style presentations and 9 for presentation in a poster session.
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Foreword

The Workshop on ML and its Applications was held on June 20 and 21, 1992, in San Francisco, California, in conjunction with the ACM SIGPLAN ’92 Conferences and Workshops.

The ML programming language has been an important tool and framework for research in language design and implementation. As the language and its implementations have matured, the range of applications has increased greatly. New applications, combined with new ideas in design and implementation, have stimulated a large number of significant activities in research and software development. This workshop, like the previous workshops in Princeton, Edinburgh, and Pittsburgh, provided a forum for these activities, with a special emphasis on applications of the language.

A total of 21 abstracts were selected for presentation and discussion at the workshop. Of these, 12 were chosen for conference-style presentations and 9 for presentation in a poster session. Four-page abstracts were submitted, almost all by electronic mail. The abstracts were not formally refereed, but were reviewed by all members of the program committee. Technical excellence, originality, and relevance to the theme of “applications” were the main selection criteria. Most of the abstracts that focused primarily on a particular application of ML, as opposed to a language extension or implementation, were grouped into the poster session. Unfortunately, many very good abstracts were not selected, in part to keep a good workshop focus, and also to leave enough room in the schedule for informal discussions. A goal of the program committee was to encourage participation by as many people as possible, while still retaining a productive workshop atmosphere.

We would like to thank all of the authors who submitted abstracts, whether accepted or not. The large number of submissions is a testament to the importance and excitement surrounding the ML language.
ACM SIGPLAN Workshop on ML and its Applications
San Francisco, California
June 20–21, 1992

Saturday, June 20, 1992

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Abstract Value Constructors

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1 Introduction

Standard ML (SML) has been used to implement a wide variety of large systems, such as compilers, theorem provers and graphics libraries; even operating systems have been contemplated. While SML provides a high-level notation for programming large applications, there are some missing language features. One such feature is a general mechanism for assigning symbolic names to constant values. We present a simple extension of SML that corrects this deficiency in a way that fits naturally with the semantics of SML [MTH90, MT91]. Our proposal generalizes SML’s datatype constructors: constants generalize nullary datatype constructors (like nil), and templates generalize non-nullary datatype constructors (like ::). Constants are identifiers bound to fixed values, and templates are identifiers bound to structured values with labeled holes. Templates are useful because they allow abstract access to user defined types. In the remainder of the paper, we give examples of the utility of our mechanism, informally explain its semantics, discuss its implementation, describe the interaction between our mechanism and the module system, and discuss related work.

2 ML datatypes and patterns

The SML datatype mechanism provides a high-level mechanism for declaring new structured types. For example, the declaration

\[
\text{datatype 'a list = nil | :: of ('a * 'a list)}
\]

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defines the standard predefined polymorphic list type. This declaration defines two data constructors: nil is a nullary constructor representing the empty list, and :: (which is an infix operator) is the list constructor. The expression 1::2::3::nil evaluates to an integer list of three elements (SML provides the derived notation [1, 2, 3] for this construction). Datatypes can also be used to define enumerations; for example,

```sml
datatype direction_t = NORTH | SOUTH | EAST | WEST
```

The utility of datatypes is greatly enhanced by the use of pattern matching in function definitions to do case analysis and value decomposition. For example, the following function takes a list of strings and returns a list with commas inserted between adjacent elements:

```sml
fun commas [] = []
  | commas [s] = [s]
  | commas (s::r) = s :: "", :: (commas r)
```

This function consists of three rules (or equations). The left side of a rule is a pattern, the right side is an expression to be evaluated when the pattern is matched. The first rule in commas matches the empty list. The second matches the singleton list and binds the list's element to s. Theoretically, the third matches any list of one or more elements, but, since the order of equations defines their precedence, it in fact matches only lists of two or more elements; it binds s to the head of the list and r to the tail. Patterns in SML are linear, i.e., a variable may occur at most once in a pattern.

The data constructors defined by datatypes have a dual nature; they are used both to construct values (when they used in expressions), and to destruct values (when they are used in patterns). We use the term value constructor to refer to identifiers with this dual nature. If no confusion with type constructors is possible, we abbreviate this to constructors. Our extension of SML provides another mechanism through which identifiers with this dual nature may be defined. Since the values constructed using the data constructors of a datatype belong to the datatype's type, they are effectively distinct from any pre-existing values. Our mechanism allows the definition of constants, that is value constructors that refer to fixed pre-existing values, and the definition of templates, that is value constructors that refer to fixed, structured values with named holes. Templates can be thought of as the constructors for families of pre-existing values in the same way that non-nullary data constructors may be thought of as constructors for families of new values.

### 3 Constants

The use of symbolic names to refer to constant values is an essential tool in the writing of understandable, maintainable software. Every systems programming language provides some sort of support for symbolic constants: C has a powerful macro pre-processor, Modula-3 has constant declarations, and so forth. In SML, val bindings allow values to be given symbolic names, but these names cannot be matched against in patterns. This is a serious
limitation because pattern matching is the principal mechanism for case analysis and the
decomposition of structured values. It is also possible, using the datatype declaration, to
declare identifiers that may be matched against in patterns; however, these are not sym-
bolic names for existing values, but rather names for entirely new values. While these two
mechanisms are adequate for many applications, sometimes what is required is a mechanism
that allows identifiers both to be bound to particular values and to be matched against in
patterns.

For example, consider the implementation of an X Window System library. The X Win-
dow System uses a device independent representation of keyboard keys called keysyms. We
need to provide users with a symbolic name for each of them. Clearly, users need to be able
to pattern match against these names so that they can conveniently write programs that
respond to different keystrokes differently. Thus val bindings are not suitable to provide
these names. Using a datatype to represent keysyms is also unsuitable. The wire represen-
tation of a keysym, that is, the representation used by the X server, is a 29 bit integer. If the
library uses a datatype to represent keysyms, it must also include a function to convert from
wire representations to library representations, and another to convert library representa-
tions back into wire representations. It is essential that these functions be mutually inverse.
Since there are literally thousands of keysyms defined — even minimum English language
support requires 512 — these functions present a maintenance nightmare. Furthermore,
the use of such huge functions adds significant run-time overhead to the marshaling and
unmarshaling of messages. Using our mechanism, we can write definitions such as

datatype keysym_t = KEYSYM of int
  const KS_a = KEYSYM 97
  const KS_b = KEYSYM 98

Here the representation of a keysym is the constructor KEYSYM wrapped around the wire
representation. (This is a standard idiom for creating new types isomorphic to existing ones.
Compilers can represent keysym_t and int identically, and treat the constructor KEYSYM
as a no-op.) The identifiers KS_a and KS_b are declared as constants that represent the
characters 'a' and 'b'. They can be used both in patterns and in expressions. The identifier
KS_a stands for the structured value (KEYSYM 97). Note that this representation of keysyms
has a further advantage over the datatype representation in that it allows keysyms from
different keysets to be defined in different modules, enabling users to import only those
keysets they actually need.

This use of our mechanism is reminiscent of the define mechanism of the C language
[KR88] (arguably, this is one area in which C provides a higher-level notation than SML).
SML has a more general pattern matching facility than the C switch statement, and our
symbolic constants reflect that. For example, a calendar program might include the follow-
ing definitions

datatype date = DATE of {month : int, day : int}
  const CHRISTMAS = DATE {month=12, day=25}
4 Templates

Templates are a natural generalization of symbolic constants to allow labeled holes\(^1\). They provide a mechanism to define a concise syntax for a collection of similarly structured values. A template is defined by a declaration of the form

\[ \text{const id trivpat} \equiv \text{patexp} \]

where trivpat is a pattern that involves only variables and record construction, and patexp is a pattern that contains only variables, record construction, special constants and other value constructors. Every patexp can be viewed both as a pattern and as an expression (they are the syntactic intersection of patterns and expressions). Every variable appearing in trivpat must also appear exactly once in patexp.

For example, the template mechanism allows us to define a template for the days of the month July using the declaration

\[ \text{const JULY(x)} \equiv \text{DATE\{month=7,day=x\}} \]

This would allow expressions like JULY(17) to be used to create date values for days in July. It would also allow code such as

```
case day
  of JULY(4) \Rightarrow "Independence Day"
  | _ \Rightarrow "not Independence Day"
```

in which JULY is used as a constructor in a pattern match.

A more substantial and more useful example of templates arises in systems that do term manipulation (such as the Nuprl proof development system, or code optimizers). For the sake of concreteness, we set our example in the context of a generalized \(\lambda\)-calculus. A term is either a variable or the application of an operator to a sequence of bound terms. For example, \(\lambda\) and \(\text{ap}\) are operators and \(\lambda(z.a)\) is a term with \(x\) bound in the sub-term \(a\), and \(\text{ap}(a;b)\) is term with sub-terms \(a\) and \(b\) (but no bound variables). One possible representation of this term language uses a different constructor for each operator.

```
datatype term = VAR of var
              | LAMBDA of (var * term)
              | AP of (term * term)
              | PLUS of (term * term)
              | ...
```

This representation has a two major deficiencies. First, functions like substitution that are independent of operators need to be written using many similar cases. Second, often

\(^1\)The term template was suggested by Dave MacQueen.
(for example, in Nuprl), it is desirable to make the set of operators extensible. But, if this representation of terms is used, the datatype needs to be changed to extend the operator set, and this requires a complete recompilation of the program. Furthermore, extension of the operator set exacerbates the problem with functions like substitution — the addition of a new operator requires that a new case be added to each such function. The following representation defines a regular, extensible structure.

```plaintext
datatype operator = OP of string
datatype term = VAR of var | TERM of operator * ((var list * term) list)
```

Unfortunately, the syntax of expressions and patterns using this representation is quite ugly. For example, the pattern that matches β-redices (i.e., terms of the form \( \text{ap}(\lambda (x.s); t) \)) is

```plaintext
TERM(OP "AP", [[(), TERM(OP "LAMBDA", [[[x], s]]), ((), t)]])
```

compared to

```plaintext
AP(LAMBDA(x, s), t)
```

in the first representation. Moreover, the second representation scheme does not provide the syntactic checking given by the first representation. \( \text{TERM}(\text{OP } \"\text{LAMBDA}\" , []) \) is a perfectly acceptable member of the type \( \text{term} \) even though terms formed with the \( \lambda \) operator should always have exactly one subterm in which exactly one variable is newly bound. Additionally, the use of strings to name operators adds the overhead of string comparison to pattern matching. All these problems are nicely solved using our mechanism. For example, with the following declarations

```plaintext
datatype operator = OP of int
datatype term = VAR of var | TERM of operator * ((var list * term) list)
```

the pattern for β-redices is again \( \text{AP}(\text{LAMBDA}(x, s), t) \). While the \( \text{term} \) type still includes many unintended values, disciplined use of the templates \( \text{AP}, \text{LAMBDA} \) and \( \text{PLUS} \) makes it impossible for users to produce these values accidentally. The required discipline can be enforced using the module facility. Thus, we get both the succinctness and syntactic checking of the first representation and the flexibility and regular structure of the second.
Constructors and the module system

The module system is an important feature of the SML language. Our proposal meshes elegantly with the module system, and the module system makes the mechanisms of our proposal even more powerful. Because the module system allows the programmer to limit the externally visible definitions of a structure, it is possible to limit the constructors available to users of the structure. For example, in the term example of the previous section, it might be desirable to limit users to the constructors AP, LAMBDA and PLUS, without giving them access to the lower level constructors such as TERM, and LAMBDA_0P. This would make it impossible for users to create junk terms.

Because data constructors are just a special kind of value constructor, it is possible to provide an interface to a structure in which they are made available as constructors without having to make their declaration as datatype constructors visible. This in turn allows the constructors of a datatype to be selectively exported. This is useful if the datatype declaration includes private constructors that are used to form intermediate values not valid in input or output.

Our proposal adds constructor specifications to the syntax of signatures. Nullary constructors are specified by the specification

```
const id : type
```

and unary constructors are specified by the specification

```
const id : type of type'
```

In this specification type' gives the type of the argument, and type gives the type of the constructed values. The syntactic distinction between unary and nullary constructors is required because, in patterns, unary constructors must always appear with arguments and nullary constructors may never appear with arguments. The legality of code such as

```
signature SIG =
  sig
    type unknown
    const K : unknown
  end

functor F (A : SIG) =
  struct
    fun f A.K = 17
    | f _ = 12
  end
```

depends on K being a nullary operator. Thus it is essential that structures such as
structure S = struct
  type unknown = int -> int * int
  const K x = (x, 12)
end

not be allowed to match SIG. We do this by explicitly associating arities with constructors rather than inferring them from type information as is done in the standard semantics. This in turn requires that the arity of constructors be available in signatures.

6 Informal Semantics

We have adapted the formal semantics of SML given in [MTH90, MT91] to handle value constructors and templates. Because of space limitations, we can only sketch the important issues here; the interested reader is referred to [AR] for the details.

Clearly, our proposal requires that a value constructor identifier can denote different values in different program contexts. For example, after the declaration

\texttt{const MyNumber} = 1

the constructor \texttt{MyNumber} must denote the integer 1, while after the declaration

\texttt{const MyNumber} = 2

it must denote the integer 2. To this end, an environment is used to associate constructor identifiers with their values. Moreover, the values associated with these identifiers must provide sufficient information to encode their behavior in patterns as well as in expressions.

The semantics of constants is straightforward. We use the environment to associate each constant identifier \(c\) with its value \(v\). Nullary data constructors can be treated in this way: they are bound to themselves in the environment. When \(c\) is used in an expression, its value is \(v\). Attempts to match a value \(w\) against \(c\) succeed if \(w = v\) and fail otherwise.

The semantics of a template \(C\) is given using a pair of functions \((C_\pi, C_\pi)\): an injection and a projection. The injection is used to construct values in expressions, and the projection is used to perform the data destructuring associated with patterns. Non-nullary data constructors can be treated in this manner. Using ML-like notation, the functions corresponding to the data constructor \(\text{TERM}\) defined above are

\[
\begin{align*}
\text{TERM}_\pi &= \text{fn } x \Rightarrow \text{TERM}\, x \\
\text{TERM}_\pi &= \text{fn } (\text{TERM}\, x) \Rightarrow x \mid _\pi \Rightarrow \text{FAIL}
\end{align*}
\]

where \(\text{FAIL}\) is a special value used to denote match failure. Similarly, the functions associated with the identifier \(\text{LAMBDA}\) by the template definition given above are

\[
\begin{align*}
\text{LAMBDA}_\pi &= \text{fn } (x, t) \Rightarrow \text{TERM}(\text{LAMBDA}_\text{OP}, [[[x], t]]) \\
\text{LAMBDA}_\pi &= \text{fn } (\text{TERM}(\text{LAMBDA}_\text{OP}, [[[x], t]])) \Rightarrow (x, t) \mid _\pi \Rightarrow \text{FAIL}
\end{align*}
\]
Note the appearance of \texttt{LAMBDAP} in these functions. Since the scope of \texttt{LAMBDAP} may differ from that of \texttt{LAMBDAS} these functions must record the environment in which they were defined.

The most important point in the semantics of the module system is the appropriate definition of signature-structure matching. Informally a signature $\Sigma$ matches a structure $S$ if it is less specific: i.e., has fewer components, less polymorphism, or elides the constant nature of values. For example, the specification

\begin{verbatim}
val nil : int list
\end{verbatim}

matches the standard list constructor \texttt{nil} (which has type 'a list). The specification

\begin{verbatim}
const nil : int list
\end{verbatim}

also matches the constructor \texttt{nil}. This can be used to export a limited view of a datatype. When a structure is constrained by a signature (e.g., when used as the argument to a function), it is necessary to thin it by removing the unused components. Thinning also involves mapping constructors to values by discarding their projection functions. To facilitate the definition of structure thinning, our semantics uses separate environments to associate the injection and projection components of a constructor’s value with the constructor rather than using a single environment and associating explicit injection-projection pairs with each constructor.

### 7 Extensions

In addition to datatype declarations, SML includes two other flavors of value constructors: exception constructors and the special constructor \texttt{ref}. Treating exception constructors as value constructors is entirely straightforward. Every time an exception declaration is evaluated, a new exception name is generated, and associated with the declared exception identifier in both the value and projection environments. Treating \texttt{ref} as a value constructor is more difficult. Such a treatment of \texttt{ref} would allow it to be used in constant and template declarations. Since the semantics of the \texttt{ref} depend on the store, this would mean that the semantics of all value constructors potentially depend on the store. Furthermore, it is not entirely clear what the semantics of the constant declared by

\begin{verbatim}
const strange = ref 1
\end{verbatim}

should even be — should every use of \texttt{strange} in an expression result in a new element of storage being allocated, or should they all refer to the same address? We do not view this as a deficiency of our proposal, rather we view it as evidence that SML’s treatment of \texttt{ref} as a constructor is a mistake.
One of the asymmetries of the design of SML is that one can define an injection-only view of a constructor (using a `val` specification), but not a projection-only view. As an example of the utility of such a mechanism, consider the implementation of points in a graphics toolkit. We may want to restrict points to some sub-range of the representable values, while still using pattern matching to decompose values. This might be done as follows

```plaintext
abstype pt_t = PTREP of (int * int)

with exception PtRange

fun mkPt (a, b) =
  if (a and b are in range)
  then PTREP(a, b)
  else raise PtRange

proj PT (x,y) = PTREP (x,y)
```

where the `proj` declaration defines an identifier that can be used only in patterns. It is safe to allow wildcards on the right-hand side of projection declarations. Because our framework separates the injection and projection aspects of constructors, it is fairly straightforward to add this kind of mechanism.

### 8 Implementation

We have built a simple testbed implementation of our proposal. At compile time, template identifiers are bound to `(trivpat, patexp)` pairs (the environment component of an injection or projection closure is coded in the representation choices for the constructors). When a known template occurs in a pattern, we inline expand it. Say that `C` is bound to `(tp, pe)` and we have the pattern `p`. We symbolically apply `tp` to `p`. This yields a substitution on the variables of `tp` (which are the same as the variables occurring in `pe`). Applying the substitution to `pe` yields a new pattern, which replaces `C p`.

For a functor parameterized by a structure with constructors, there are two implementation issues: what is the representation of the abstract constructors and how are they used in patterns. To handle the first question, we use implicit structure members for the injection and projection functions. Note that these only need to be added when the structure is made abstract (i.e., by functor application), and can be generated as part of signature thinning. The injection function is an ordinary function, while the projection is a function that returns either the sub-values or FAIL. When building a decision tree, the compiler treats abstract templates as ordinary constructors, which allows merging of matches against them. A test against an abstract constructor is a call to the projection function. Of course, the use of functions to implement the construction and destruction associated with abstract constructors may result in a certain degradation of performance. Of particular concern is the loss of merging when two abstract constructors have common structure. (For example,
the templates LAMBDA and AP defined earlier share the structure \(\text{TERM(OP \_ \_ \_)}\). Any value not matching this pattern cannot possibly match either of them. Other costs include the loss of inline tests and the replacement of branch tables with trees of conditionals. Note that these costs are incurred only when constructors are actually abstract, that is, when functors are used. In particular no penalty is incurred when structures, which have transparent signatures, are used rather than functors. Moreover if macro expansion were used in the implementation of functor application (a reasonable thing to do when compiling a production version of a system) functors would reduce to structures and abstract constructors would become concrete.

Our implementation of abstract constructors can also be applied to solve an outstanding problem with datatype representation and abstraction. In some implementations of SML [App90], the representation of the datatype defined by \(\text{datatype } d = A \mid B \text{ of } \tau\), depends on the representation of \(\tau\). If the representation of \(\tau\) is appropriate, it is possible to represent the value \(B \text{ exp}\) with the representation of \(\text{exp}\). Problems arise when \(\tau\) is abstract, since its representation is known at the definition of \(d\) but not elsewhere. Extending our technique to cover abstract datatype constructors that fall into the danger zone solves this problem. Although our solution to the problem incurs a performance penalty, a less speedy program is better than one that does not run correctly.

9 Related work

Wadler's view mechanism ([Wad87]) shares the objective of allowing data abstraction and pattern matching to cohabit. Views were once part of the Haskell definition ([HW88]), but were dropped because of technical difficulties. Conceptually, a view of a type \(T\) is a datatype \(T'\) together with a pair of mappings between \(T\) and \(T'\). Ostensibly these maps are isomorphisms, but since they are defined by the user, there is no assurance that the types are truly isomorphic. Views and templates differ in several significant ways. The principal difference is that Wadler's views define maps between concrete representations, whereas templates provide abstract views of a single representation. Because views define different types, a given pattern match can involve only one view. In addition, once a view is defined, it is not possible to add additional constructors (even if other representations admit additional objects). Templates, on the other hand, do not suffer these restrictions. The implementation of views uses the user defined maps to convert between representations; thus, pattern matching can incur arbitrarily large performance penalties\(^2\). In our scheme, most uses of templates incur no run-time cost, and even in the worst case they add only a constant overhead to pattern matching. Our presentation of the semantics of templates is more detailed than that of views given in [Wad87]; furthermore, we address the semantic and implementation issues related to separate compilation and parameterized modules.

The CAML system ([WAL+]) provides a mechanism for defining new concrete syntax, by specifying a grammar to map quoted phrases to the internal representation of programs. This mechanism could be used to implement our template mechanism, although the imple-

\(^2\)In fact, there is no guarantee that the maps even terminate.
mentation details appear non-trivial. Recently, a quotation mechanism has been proposed for SML, which allows terms in some object language to be included in expressions ([Sli91]). This provides some of the syntactic convenience of our mechanism, but it provides no help for pattern matching against terms of the object language.

References


Efficient representation of extensible records

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April 10, 1992

Abstract

We describe a way of representing polymorphic extensible records in statically typed programming languages that optimizes memory allocation, access and creation, rather than polymorphic extension.

Introduction

Type systems for records have been studied extensively in recent years. New operations on records have been proposed such as polymorphic extension that builds a record from an older one without knowing its fields. Such operations are very powerful, and were not always provided as primitive constructs in untyped languages.

In comparison to the numerous results on the type theory of records, there has been less interest in their compilation. Many languages still have monomorphic operations on records, e.g. most implementations of ML [HMT90, Wei89, Ler90]. Others, that have more powerful records, use association list techniques, eventually improved by caching.

Safe untyped languages require that the presence of fields is checked before access. The use of association lists interleaves dynamic checking with access to fields. In strongly typed languages, the presence of fields is statically checked. Thus the representation of records by association lists performs superfluous runtime checks, and it seems that cheaper solutions could be found.

We propose a representation of records based on a simple perfect-hash coding of fields that allows access in constant time with only a few machine instructions which can be dropped to a single instruction whenever the set of fields of the record is statically known. Creation can be performed in time proportional to the size of the record, and allocates a vector of size the number

of fields plus one. Only the polymorphic creation of records has to pay more in time and memory.

In section 1, the specificity of records as data product structures serves as an introduction to the conditions for which our representation will work in practice. The method is described in detail in section 2 as the encoding of partial functions from labels to values with finite domains. In section 3 we extend the method to records with defaults. These are total functions from labels to values that are constant almost everywhere. As an application we get safe standard records in an untyped language. In section 4, we discuss how to handle pathological cases in order to prevent bad behavior.

1 Records and their specificity

Records are product data structures. Each piece of information is stored with a key, more commonly called a label, that is used to retrieve the information. There is at most one value associated to a label. By field a pair noted $\alpha \mapsto v$ of a label $\alpha$ and a value $v$. Such data structures are of common use in computer science. However our definition of records is too vague for choosing a good representation of records. It is necessary to know the average and the maximum size of these structures, to know whether they are created incrementally, the frequency of the different operations and which ones are privileged. It is obvious that different programs will give different answers. These questions can only be answered in general, or the answers that we give below can also be taken as assumptions for which our representation of records will work in practice.

Records are provided with three operations. Creation builds a record with a finite number of labels together with values associated to these labels. Extension takes a record and builds a new one that has all fields of the first one plus a finite number of new fields. Access takes a record and a label and returns the value associated to that label. It fails if the label is not in the domain of the record.

Records have a relatively small number of labels. At most a couple of hundred, on average less than ten. Non incremental creation and access are both very frequent and are privileged. There are usually many records with the same domain. Space and time are equally
important.

The simplest representation of records by association lists is good for very small records but it makes access to large records too slow. Balance trees would have better performances for large records but the overhead has also to be paid for small records; they also require too much memory. General hashed tables will also have an overhead that is not acceptable for small records.

2 Extensible records with polymorphic access

In this section we consider records as partial functions from labels to values, with finite domains. The problem is to find a representation for such functions, such that under the assumptions of the above section, the operations on records can be performed efficiently. By performing, we mean evaluating in general. In particular, this applies to compilation where some of the evaluation can be done statically.

Standard monomorphic operations on non extensible records should not be penalized by the introduction of more powerful records. Although it is always possible to keep two kinds of records coexisting, the new records should replace the older, weaker ones. It should be left to the compiler to recognize that some record operations are monomorphic and thus can better be compiled. Indeed, this will not be possible for all compilation schema.

A record \( r \) is a partial function from labels to values with a finite domain \((a_i \mapsto v_i)_{i \in [1,n]}\). The simplest decomposition of this function is

\[
\begin{align*}
\text{Labels} & \quad \mapsto \quad [1,n] \\
\mapsto \quad \text{Values}
\end{align*}
\]

\( a_i \mapsto i \mapsto v_i \)

The total function \( v \) stores the components of the record, while the the partial function \( h \), called the header, describes how labels are mapped to indices. This decomposition suggests the representation of \( r \) in a vector \( H \):

\[
\begin{cases}
0 \mapsto H \\
i \mapsto v_i & i \in [1,n]
\end{cases}
\]

where \( H \) represents the function \( h \).

The partial function \( h \) needs to be defined at least on the domain of \( r \) and it should better be injective on the domain of \( r \) too. Any such function would work, since \( v \) is then defined by

\[
(h \circ \text{dom}(r))^{-1} \circ r
\]

up to permutation of indexes with identical values.

If all records are coded such that their headers (the representations may differ provided they implement the same function) only depend on their domains, then compilation of operations on records whose set of fields is statically known can be optimized by partially evaluating their header. The access become a single indirect read to fetch the value of the record on that field. The creation is always in that case, since it builds a record with \( n \) fields from nothing. The header can be computed statically and shared between all records built by the same function. The cost is reduced to allocating and filling \( n + 1 \) fields of a vector. More generally, headers can be shared between all records that have the same domain by keeping all existing headers in a table.

Polymorphic access and polymorphic extension must use the headers. For sake of simplicity, we consider that labels are integers. The parser and the printer would deal with the isomorphism between integers and names in a real language.

Finding a good representation of \( h \) seems as difficult as finding a good representation of \( r \). There are two differences, though. Since the header is shared, we are allowed a little more flexibility on the size of \( H \). Also, \( h \) is a function on integers, thus we are allowed to use arithmetic and logic operation on integers. There is no hope of finding a direct representation of \( h \) by arithmetic operations, since its domain is completely arbitrary. At least some mapping between integers has to be an arbitrary map represented by a vector of integers for instance. A mixed decomposition of the header \( h \) is:

\[
\begin{array}{c}
0 \mapsto (\mod p) \\
[0,p-1] \quad \eta & \quad [1,n] \\
\end{array}
\]

where \((\mod p)\) is injective on the domain of \( r \). Such a decomposition is always possible, to the price of a higher \( p \). In practice the smallest \( p \) that works is on average twice the size of \( n \) for a few labels and three to four times for larger sets of labels. Since the header is shared, this is very acceptable.

The interest of this decomposition of \( h \) is that it can be compiled efficiently and coded in a vector \( H \):

\[
\begin{cases}
0 \mapsto p \\
j \mapsto \eta(j-1) & j \in [1,p] \\
\end{cases}
\]

The partial function \( \eta \) must be extended into a total function on \([1,p]\). We write it \( \tilde{\eta} \) the unconstrained extension of \( \eta \).

In too cases below, We will also be interested in two particular extensions of \( \eta \) below that we write \( \eta_0 \) and \( \eta_1 \). The former \( \eta_0 \) is an extension of \( \eta \) with values of \([1,n]\), thus it makes \( r \) a total function. The later \( \eta_1 \) extends \( \eta \) outside of \([1,n]\), for instance \( 0 \), which provides a membership test to the domain of \( \eta \) by testing \( \eta_1 \) for equality to zero.

The domain of \( r \) must also be coded in \( H \) for polymorphic extension. All its labels can be listed at the end of \( H \):

\[
\begin{cases}
0 \mapsto p \\
1 \mapsto \eta \\
2 + j \mapsto \eta_0(j) & j \in [0,p-1] \\
1 + p + i \mapsto a_i & i \in [1,n]
\end{cases}
\]
The access can be optimized whenever the domain of the record, and consequently the header, are statically known. Such information is expected to be found by the typechecker. This cannot always be the case, however.

In order to rely on the types to know the domains of records, the attendances of fields, given by the types of records, must correspond exactly to their domains. This implies that the restriction of a record on a field modifies its header, since it changes the attendance. This is one possible semantics for restriction. Another one is to take the restriction of fields as a retyping function, that is, a function that evaluates as the identity. The choice is between an expensive active restriction that allows access optimizations or a cheap retyping restrictions that forbids them.

3 Records with defaults

In this section we consider records as partial functions from labels to values, constant almost everywhere. The problem is now to recognize whenever the field does not belong to the domain of the record (we mean the explicitly defined values, here), in which case the default value is returned. The membership test might be expensive in time or in space.

There is a cheap solution based on the same technique as above. In fact, we coded records by total function on labels, and described the domain separately in order to implement the extension of fields. Thus we could apply them outside of their domain (but get a value of unpredictable type).

Let \( r \) be a record. Consider the record \( r' \) equal to \( \text{id} \restriction \text{dom}(r) \). An arbitrary label \( a \) is in the domain of \( r \) if and only if it is equal to \( r'.a \). The auxiliary record \( r' \) only depends on the domain of \( r \) is already be coded in the header \( H \) as the domain of \( r \). Remember that \( i \) is the index in \( R \) where the value of label \( a_i \) is stored. Thus it is the index where \( a_i \) is in \( R' \), which is also in \( H \) at position \( 2 + p + i \), provided \( i = \eta(a_i \mod p) \).

We simply shift the indices in \( R \) to place the default value at position 1:

\[
\begin{align*}
0 & \mapsto H \\
1 & \mapsto a \\
1 + i & \mapsto v_i, \ i \in [1, n]
\end{align*}
\]

We compute the application of \( r \) to the label \( a \) as follows. First compute the index \( i \) associated to \( a, \) that is \( H.(a \mod (H.0)) \). If \( H.(2 + p + i) \) is equal to \( a \), then the label belongs to the domain of \( r \) and the result is \( R.(1 + i) \) otherwise it is the default \( R.(1) \).

One must be careful to use the \( \eta \) extension of \( \eta \). The \( \eta \) extension is still possible, but the above membership test must be preceded by a membership test to the domain of \( \eta \), and in case of failure the default value should also be used.

In fact, the encoding of records with defaults can be easily adapted to implement safe classical records in an untyped language: a dynamic type error is raised when the membership test fails instead of returning the default.

4 About efficiency

This section does not consider the case of records with defaults for sake of simplicity, but it can be adapted very easily to them.

The previous representation of records is very attractive since it implements linear time access, uses very little memory, keeps the same performance on monomorphic records as if all records were monomorphic. However we must check the following points. First, that the size of the header does not get too large. Secondly, that the polymorphic extension does not have too bad performance, even though it is not privileged. Last, that pathological cases can be handled.

The computation of the integer \( p \) is at the heart of every question. The problem is given a set of integers \( D \), find a small integer \( p \) such that \( (\_ \mod n) \) is injective on \( D \). The integer \( p \) does not need to be minimal, even if computed at compile time, since a larger header might make polymorphic extension more efficient. However, the minimal \( p \) gives a lower bound on the size of the header. For instance if \( D \) is randomly chosen, and the set of labels is large enough in comparison to the size \( n \) of \( D \), the probability that \( p \) disambiguates \( n \) integers is

\[
\frac{1}{(p - b)! \, p^n}
\]

The formula that gives the average smallest \( p \) in function of \( n \) is simple, but figure 1 gives an experimental result on the average \( p_{ave} \) and the largest \( p_{max} \) for on a hundred runs per column.

For small records, \( 10^4 \) runs did not give very different maximal \( p \). The dispersion of \( p \) is shown by figure 2.

The figures show that under 30 labels, the size of \( D \) does not exceeds, in average, four times the number of fields, and exceeds very rarely twice the average. For large records (above 50 fields) the header becomes very large. It is clear that another solution must be applied for large records. Even if one wished to push this limit to a hundred labels, there is always a rank that can be reached in practice (even if it is pathological) for another representation should be used.
Since, we cannot avoid a mixed representation, if we want to handle large size records in order to represent them with reasonable size headers, we use tags to distinguish between the two representations. For instance, negative integers can be used as tags for an alternate representation. There are many possibilities for the alternate representations, and since these are pathological cases, in the sense that they do not meet the requirement that we set in section 1, we do not care much about the efficiency of the alternate representation. We propose, two possible solutions that fit well with the regular representation.

The first solution is whenever \( a \) and \( b \) are equal to \( j \) modulo \( p \) to assign \( \eta(j) \) with an integer \( -q \) such that \( q \) is in \([2, p]\) and modulo \( q \) distinguishes \( a \) and \( b \) and with values that are not in the image of \( \eta \).

The second solution replaces perfect hashing by hashing with linear probing ([Sed88], Chapter 16). The header is \( H \) is

\[
\begin{align*}
0 & \rightarrow p \\
1 & \rightarrow n \\
2 + j & \rightarrow \eta(j) & j \in [0, p - 1] \\
1 + p + i & \rightarrow a_i & i \in [1, n] \\
2 + n + p + j & \rightarrow a_{\eta(j)} & j \in [0, p - 1]
\end{align*}
\]

for labels that do not conflict. When \( 2 + n + p + (a, \text{mod} p) \) is already occupied, the label \( a_i \) is placed at the smallest free position after, say \( j \), and \( \eta(j) \) is filled with \( i \).

The first method still gives access in linear time, but headers are more difficult to compute. The may be much faster on average, but require larger headers. In both cases there is a lot of freedom on how to chose \( p \), according to how many conflicts are accepted. Letting \( p \) be about 3 times the size of \( r \) may avoid searching for optima while limiting the number of clashes. The first method is more flexible, since a conflict for one label does not need to double the size of the header or recompute another header: it simply uses the holes of the actual header. Then, it can also be used even for average size records in some cases in order to compile more efficient extension.

The efficiency of polymorphic extension has not been considered yet, since it was not a privileged operation. It has to dynamically compute a new header, which may be very expensive — and at least proportional to the size of the record it extends. The time for computing the header now becomes important. There are different cases (we exclude large size records, that should be represented otherwise):

- There is no need to compute a new \( p \),
- The average case for computing a new \( p \),
- The worse cases for computing a new \( p \).

In the average case, computing a new header means that 3 different \( p \)'s must be probed per extension, since headers are in average 3 times the size of \( n \). The optimistic unit cost \( U \) is the one of a loop that contains at least one vector write and one modulo instruction. The cost for a probe is \( pU \), since the failure is probable to happen at the end. Thus the average cost for creating a new header is \( 3pU \) plus two other \( pU \) for filling the header.

However, a record with a very compact header may be extended with a label that will make the header get closer to its average size. Then about \( nU \) probes may be needed, making the cost for the new header increase to \( pnU \).

On the other hand, it is very probable that the extended header already exists or is trivial. If the label \( a \) of the extended field is taken at random, there is a probability \( (p - n)/p \) that the \( n + 1 \) fields will still be disambiguated by \( p \). In that case, the cost for creating a new header is the same as copying the old one plus modifying a few fields \( pU \).

The creation of a new header may be avoided most of the time since it is very probable that it has already been created. This requires that all existing headers are stored in a dictionary. This will save both space and time. The keys in the dictionary are the domains of headers that can be kept ordered in \( H \), so that equality tests are not too expensive, and the whole cost of searching would be lower than the minimal cost of extension.

Active restriction of fields can be implementing along the same ideas. The header is looked up in the table. If it does not exist, then the new header need not be the smallest one, provided that it is of reasonable size. This avoids the expensive cost of finding the smallest integer modulo which all elements of the domain are distinguished.

5 Other compilation schemas

There are three different ideas in the above representation of records

1. The value of fields and the position of fields are represented separately. The header that describes the later is shared between all records having the
same set of fields, two records with the same domain always have the same fields at the same position.

2. The header can be represented by a modulo followed by a projection.

3. Different representation of the headers can live together.

The first point is crucial in our representation. Any representation of records that does not originally respect this point can still be used to implement headers, then values can be stored in vectors as above. Sharing of headers will save the large amount of space required by association lists or balanced trees.

The representation of the header itself is not important. We described one possibility that is very convenient for small and medium size records. But many other representations are possible. We choose to represent the header as a structure that is interpreted both by extension and access. It could also be a closure for the access part, together with a description of the domain that is needed for the extension. This is the tag vs closure duality.

If the extension and the restriction of fields are themselves closed with the header, records could really be viewed as objects with two methods for access and extension.

Conclusion

We have presented a way of representing records with or without defaults that allows efficient access and creation. Only the more powerful features such as polymorphic extension, or true restriction of fields have to pay a higher price.

Our representation of record with defaults can be used to implement safe access in an untyped language.

An orthogonal application could be the representation of feature terms that are very related to records.

Thanks

These ideas originate in discussions with Xavier Leroy. They have been mentioned for the first time in [Ler90] and tested in the untyped version of Zinc, the ancestor of Caml-Light.

References


[Ler90] Xavier Leroy. The ZINC experiment: an economical implementation of the ML language.
A General and Practical Approach to Concrete Syntax Objects within ML†

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Abstract: In this paper we present an approach to concrete syntax object within ML, which is both general and efficiently implementable. These language enhancements add BNF rules for abstract syntax declarations and “semantic brackets” [I ... l] with inline concrete syntax and pattern matching for readability and conciseness. This approach has several improvements integrated together which either do not appear in previous works, or appear in forms which are very restrictive or have very inefficient implementations. Our improvements are: (1) inline concrete syntax within “semantic brackets” has been integrated both with the ML type system and the ML scope rules, (2) concrete syntax can appear both as syntactic patterns for pattern matching and as syntactic expressions for building objects, (3) patterns can be nested to arbitrary depth, (4) concrete syntax and ML objects can be mixed; so called anti-quotations are supported directly, (5) patterns and parts of patterns can be augmented with type information, (6) efficient integration with a general incremental LR(1) parsing mechanism.

These extensions have been efficiently implemented within our DML system. DML, the Denotational Meta-Language, is a dialect of Standard ML with extension aimed at making it (even more) useful for implementing denotational specifications of programming languages.

1. Introduction

Functional languages generally have some facilities for defining and computing with inductively defined data types. Unfortunately, only the simplest prefix syntax is available for objects of these types. It is generally true for problem domains where data objects with syntactic structure are specified and later accessed and manipulated, i.e. when a meta-language computes with an object-language, that the implementation tends to be cluttered with prefix constructors and/or access functions for abstract syntax and other intermediate forms. Typical examples are conventional compilers and implementations of denotational specifications of programming language semantics. If context-free grammatical methods can be used to specify this structure, then a set of language extensions based on inline concrete syntax, e.g. as proposed in this paper, can lead to more compact and readable code. The language extensions described in this paper, implemented within our DML system, have so far been used primarily in compiler generation from denotational semantics specifications. An overview of the system is given in [PF92], while this paper concentrates on the

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implementation details of the concrete syntax extensions.

2. Related Work

The work by Aasa et al [APS88] describes an approach to inline syntax, which in its current form only applies to sequences of characters and is so inefficient that it is not practically usable. Its inefficiency is mainly due to its use of Earley's [Earley70] algorithm which repeatedly recomputes parsing automatons which are reused in our method. An implementation has been done, hidden within the LML (Lazy ML) system, but it is not used by LML itself [Augustsson91].

Lee [Lee89] has a very simple but restrictive approach to implementing in-line syntax within semantic brackets in the context of denotational semantics specifications. A syntactic pattern is converted to a string by simply concatenating everything between the semantic brackets. A similar conversion occurs for each production in the syntax declaration. Patterns in semantic equations are then resolved by searching linearly for matching strings from the set of productions. This disallows patterns with different variable names but similar structure, and patterns with nested structure. These restrictions are not present in the DML system.

The Refine language [Refine90] contains perhaps the most practical and complete implementation of inline concrete syntax before our work, but still lacks integration with a general type inference mechanism and often has problems with parsing conflicts due to its use of standard LALR(1) techniques.

The CAML system from INRIA [Weis et al 89] also includes a facility that can be used to provide concrete syntax for datatypes. In CAML, a grammar declaration consists of a set of productions, each associated with an ML expression that builds the appropriate resulting term. The grammar is sent to Yacc, and the parsing tables are read back in and mapped to a set of parsing functions. These functions can be applied to strings, files or the current input. When parsing, the input is pre-tokenized by the ML system into a stream of identifiers, strings and numerical constants. It is possible to mix concrete syntax and ML expressions (so called anti-quotations) since the system's parser for ML expression is visible as the function parse_caml_expr, and can be called from the user's code in the grammar rules. The special syntax "<<...>>" is used to delimit parts of an ML program that should be parsed according to user-defined grammar. There is a notion of the current grammar which is used by default: <:grammar:non-terminal<...>> can be used to explicitly direct the focus to a particular grammar and (optionally) a particular start symbol.

The system appears to be most suitable for providing a simple means to prototype parsers for the concrete syntax of programs when developing compilers or interpreters. Since it runs entirely before the type-inference process, no ambiguities can be allowed, even if type constraints would resolve them. Overall, the system has a more "programmatic" than declarative feel. While our approach doesn't support the building of user-level parsing functions, we think it is better integrated with the ML language, especially since it uses type information in addition to context-free syntax, and from the start supports anti-quotations without the need for extra programming.
3. The Use of Syntax Definitions in Denotational Semantics

A common way to specify the type of syntax trees in texts on denotational semantics is to use BNF rules like this:

\[
\begin{align*}
C : & \text{Con} \\
I : & \text{Ide} \\
E : & \text{Exp} \\
E ::= & C \mid I \mid E + E
\end{align*}
\]

and then refer to syntax tree objects by writing their syntax between "semantic brackets":

\[
\begin{align*}
eval[C] & \text{env} = C \\
eval[I] & \text{env} = \text{env} I \\
eval[E_1 + E_2] & \text{env} = (\eval[E_1]\text{env}) + (\eval[E_2]\text{env})
\end{align*}
\]

The definition-by-syntax facility is really just a shorthand, eliminating the need to explicitly deal with Cartesian products, disjoint unions and their appropriate injections, projections and tag tests. In DML we take the same view: syntactically defined types provide a convenient notation for ordinary types defined with datatype declarations. The example could in DML be expressed as follows:

\[
\begin{align*}
type & \text{Con} = \ldots \\
type & \text{Ide} = \ldots \\
syntax & \text{Exp} = \text{Con} \mid \text{Ide} \mid \text{Exp} "\text{+}" \text{Exp}
\end{align*}
\]

\[
\begin{align*}
\text{fun eval} & [\text{Con}] \text{env} = c \\
\mid & \text{eval} [\text{Ide}] \text{env} = \text{env} i \\
\mid & \text{eval} [\text{Exp} "\text{+}" \text{Exp}] = (\eval[\text{Exp}]\text{env}) + (\eval[\text{Exp}]\text{env})
\end{align*}
\]

Translating it to vanilla SML might result in:

\[
\begin{align*}
\text{datatype exp} & = \text{ConExp of Con} \\
\mid & \text{IdeExp of Ide} \\
\mid & \text{AddExp of Exp \ast Exp}
\end{align*}
\]

\[
\begin{align*}
\text{fun eval} & (\text{ConExp c}) \text{env} = c \\
\mid & \text{eval} (\text{IdeExp i}) \text{env} = \text{env} i \\
\mid & \text{eval} (\text{AddExp(\text{Exp}, \text{Exp})}) \text{env} = (\eval[\text{Exp}]\text{env}) + (\eval[\text{Exp}]\text{env})
\end{align*}
\]

The transformation is non-trivial in general. In this example, the "pure" SML version wasn't too unreadable, but more complex types and patterns (especially nested ones) are clearly more conveniently handled using grammar rules. There are two basic reasons for preferring syntactic type definitions. First, we do not have to cast all expressions and patterns into the prefix or binary infix syntax of SML. Instead we can use (almost) any meaningful notation we wish. Second, grammars allow us to make "unit productions" (type conversions) implicit. Consider:
Here DML enables us to make the mapping from integer constants to Exp:s implicit, whereas SML wants us to spell out all the gory details.

4. The implementation of DML's syntactic objects

This section describes in detail how the extensions for syntactic objects are implemented. First the facility itself is preserved, and an example is given to show that syntax alone cannot be used to resolve syntactic objects; types are needed as well. Then the compiling strategy is outlined, followed by detailed descriptions on how the pseudo-parallel LR(0) automatons used here are defined and implemented.

4.1 Grammar of the DML syntax extensions

Table 1 contains the grammar for the DML syntax extensions in the same style as the Standard ML definition [HMT90]. We use string to denote the class of all string constants, whereas the definition of Standard ML conflates all constants to the class scon. <thing> means that thing is optional.

| dec       | := | syntax synbind <withtype typbind> |
| synbind   | := | tyvarseq tycon = srules <and synbind> |
| srules    | := | <srule> <srules> |
| srule     | := | string <srule> |
|           |   | ty <srule> |
| atexp     | := | [l <synexp> l] |
| synexp    | := | string <synexp> |
|           |   | atexp <synexp> |
| atpat     | := | [l <synpat> l] |
| synpat    | := | string <synpat> |
|           |   | atpat <synpat> |

4.2 Syntactic ambiguities

Purely syntactic preprocessing is often not enough to determine the constructor form of a syntactic expression or pattern. Syntactically ambiguous patterns are common in denotational semantics specifications.
There, implicitly-typed variables are used to disambiguate them.

**Syntactic Domains:**

\[
\begin{align*}
N & \in \text{Nml} & \text{(numerals)} \\
I & \in \text{Ide} & \text{(identifiers)} \\
E & \in \text{Exp} & \text{(expressions)}
\end{align*}
\]

**Abstract Syntax:**

\[
\begin{align*}
E & ::= N | I | E_1 + E_2 \\
E & : \text{Exp} \rightarrow \text{Env} \rightarrow \text{Int} \\
N & : \text{Nml} \rightarrow \text{Int} \\
E[N]_\rho & = N[N] \\
E[I]_\rho & = \rho[I] \\
E[E_1 + E_2]_\rho & = E[E_1] + E[E_2]
\end{align*}
\]

Here the implicit typing of \(N\) and \(I\) act to disambiguate the first two patterns. In this work, we wanted for syntactic patterns to behave as much as possible like ordinary SML patterns without having to introduce additional declarations or limit the way patterns may be formed (as in Lee’s MESS system).

DML would in the first equation find that the type for \(N\) had been constrained to be a numeral, and in the second equation that \(I\) had to be an identifier. For this example this is enough to completely resolve the apparent ambiguities. In those cases where the implicit type constraints aren’t enough, the ordinary SML \texttt{typ} construct can be used to explicitly add the needed constraints (much like how overloaded variables sometimes need this). Note also that this problem is largely eliminated with the use of “noise” keywords in syntax productions.

### 4.3 General Implementation Strategy

1. Syntax is seen as a grammatical alternative to ordinary datatype declarations. In the same vain, we see syntactic expressions and patterns as alternatives to ordinary expressions and patterns built using constructor applications.

2. Every syntax declaration is implicitly transformed to a datatype declaration, derived as follows:

Given a production rule \texttt{srule}, its constructor name is built by appending the name of the non-terminal (the \texttt{tycon}), a string "::=" and the names of the items in the right-hand side. The name of a terminal is itself, the name of a type is ".". For the same rule \texttt{srule}, the arity of the constructor is determined by building a tuple type \(\texttt{ty}\) of the embedded types (not the terminals) in the right-hand side. If \(\texttt{ty}\) is the empty tuple type, then the constructor is a constant, otherwise it is a function with domain \(\texttt{ty}\). For example:

\[
syntax \texttt{Stmt} ::= \texttt{"if" Exp \texttt{"then"} } \mid \texttt{"else" Stmt } \mid \texttt{"skip"}
\]
has the following corresponding datatype declaration:

```plaintext
datatype Stmt = "Stmt::=if_then_else_" of Exp*Stmt*Stmt
 | "Stmt::=skip"
```

Just as for datatype declarations, it is an error for a synbind to bind the same tycon or (generated) constructor more than once.

3. Occurrences of syntactic objects are translated to ordinary objects (expressions or patterns) built using explicit constructor applications. For every syntax declaration, its productions are added to an incrementally-built pseudo-parallel LR(0) parser[1]. Each syntactic object is first parsed by the pseudo-parallel LR(0) automaton which results in a set of possible parse trees. When all surrounding type constraints are known (at the end of the type-checking pass) only those trees that are consistent with later type assumptions are considered. If only one such tree remains, then that is taken as the meaning of the syntactic object.

4. Given that syntax declarations has had their corresponding datatype declarations determined, and that all synexps and synpats has been rewritten to ordinary expressions and patterns, we see that no extra constructs remain for the code generator. The run-time costs of these syntactic constructs is therefore the same as if ordinary datatypes had been used.

4.4 Resolving Syntactic Objects
This section explains how syntactic objects are parsed and rewritten to ordinary SML constructs. This process is integrated with the ordinary type checking process since the two are interdependent.

The static context \( C \) is extended with a grammatical environment \( \text{GrmEnv} \). The idea is that a grammar is a finite mapping from constructed types (\( \text{ConsTypes} \)) to finite sets of productions, where each production is a sequence of terminals (string constants) and non-terminals (types). Grammatical environments follow the same scope rules as the type environments from datatype declarations.

\[
\text{GrmEnv} = \text{ConsType} \rightarrow \text{setof(Prod)} \\
\text{Prod} = \text{Token*} \\
\text{Token} = \text{String} \cup \text{Type}
\]

4.4.1 LR(0) construction
Given a context \( C \), an LR(0) parser for the grammar environment in \( C \) is constructed as follows. First the LR(0) sets of items are generated for the grammar in \( C \), where every bound ConsType is a valid start "symbol". This is achieved by introducing a dummy start symbol \( S' \), and adding the production \( S' \rightarrow \text{ty} \), for every bound ConsType \( \text{ty} \). Non-grammatical types are effectively treated as terminals.

[1] These machines have the recognizing power of LR(1).
Then the transitions of the parser are defined by the following rules, which are based on the standard ones in [HU79][2]. Note that the rules do not disallow shift/reduce or reduce/reduce conflicts. We define a parser configuration as a 4-tuple \((pstk, astk, \sigma, \text{input})\). This is just the normal LR-parsing configuration extended with a result stack and a type substitution.

- \(pstk\) is a stack of state numbers (the usual parse stack)
- \(astk\) is a stack of abstract syntax trees (the “result” stack)
- \(\sigma\) is a finite substitution from type variables to types
- \(\text{input}\) is the remaining sequence of input tokens, terminated by the special terminal $$. Each token is either a terminal string, or a pair \((ry, x)\) where \(x\) is an ML syntax object (an \text{Exp} or a \text{Pat}), and \(ty\) is its type.

**INIT:**

\[\begin{aligned}
(q_0, [], \emptyset, \text{input})
\end{aligned}\]

Init is the initial configuration, where \(q_0\) is the start state of the automaton.

**ACCEPT:**

\[\begin{aligned}
(q, q_0, [ast], \sigma, [\$])
\end{aligned}\]

Accept is an accepting configuration if \(q\) contains a complete item \(S' \rightarrow ty^*\). The result is \(ast\).

**SHIFT:**

\[\begin{aligned}
(q::qs, astk, \sigma, str::\text{input}) \Rightarrow (q'::qs, astk, \sigma, input)
\end{aligned}\]

Shift if there is a transition labeled by the terminal \(str\) from \(q\) to \(q'\).

**GOTO:**

\[\begin{aligned}
(q::qs, astk, \sigma, (ty;x)::\text{input}) \Rightarrow (q'::qs, \text{LEAF}(\sigma'(ty),x)::astk, \sigma'*, \text{input})
\end{aligned}\]

Goto if there is a transition labeled by some type \(ty'\) from \(q\) to \(q'\), \(\text{Unify}(\sigma(ty),\sigma(ty'))\) succeeds with most general unifier \(\sigma'\), and \(\sigma'\) is the composition of \(\sigma\) and \(\sigma'\).

**REDUCE:**

\[\begin{aligned}
(q_1::\cdots::q_k::qs, ast_1::\cdots::ast_n::astk, \sigma, \text{input}) \Rightarrow (q'::qs, \text{NODE}(\sigma(ty),con,[a_1, \cdots, a_n])::astk, \sigma, \text{input})
\end{aligned}\]

Reduce if \(q_1\) contains a complete item \(ty \rightarrow prod\), \(\text{length}(\text{prod}) = k\), the number of types in \(\text{prod}\) (its arity) is \(n (n < k)\), and \(\text{con}\) is the name of the constructor for this production.

The resulting parse trees are described by the following type:

```
datatype 'a Ast = LEAF of Type * 'a
  | NODE of Type * Con * 'a Ast list
```

Leaf nodes correspond to inserted ML objects (anti-quotation), while internal nodes correspond to the application of some (syntax) constructor to a sequence of sub-trees. Each node is also tagged with its type. The type parameter \(\text{'a}\) is the compiler’s type of the leaf ML objects, corresponding to expressions or patterns.

### 4.4.2 Modifications to the Basic Type-Checker

The DML type-checker consists of two major components: the ordinary type-checker for the SML subset, and the new combined parser/type-checker for syntactic objects. The type-checker for the SML subset is implemented by combining the general structure and rules of the formal definition of Standard ML. [2] Actually, what we are constructing is a viable-prefix recognizing NFA for the LR(0) items.
[HMT90], and an imperative implementation of type unification and treatment of polymorphic types in [Cardelli87]. It is organized in two passes: Pass 1 is the ordinary top-down traversal which maintains type environments and checks type compatibility by unification. Occurrences of flexible record patterns and overloaded variables are remembered in a separate set of cleanup actions, since they in general aren’t fully determined at that point in the traversal. Pass 2 finally iterates through the set and checks that all flexible record patterns and overloaded variables have been fully determined by type assumptions made later in Pass 1.

To deal with syntactic objects, the following extensions were made:

Pass 1 calls the syntax parser every time a syntactic object is encountered. If the resulting parse forest is ambiguous, then the occurrence is remembered in the set of cleanup actions and the syntactic object is assumed to have an unconstrained type $\tau$. Otherwise, the unambiguous result is patched into the ML syntax tree directly and its type is used as the type of the syntactic object.

Pass 2 tries to disambiguate remembered parse forests by a filtering process: any parse tree whose type is inconsistent with type assumptions made later in Pass 1 is removed. If only one tree remains, it is patched into the ML syntax tree. Otherwise, there is an error (ambiguity or inconsistency).

4.4.3 Incremental generation of the automata

Since constructing the LR(0) automaton is a costly process, we do not want to waste time by redoing this each and every time a syntactic object needs to be parsed. The grammar environment in the static context is therefore accompanied by a cached, partially-built automaton that exactly mirrors the set of productions that currently are in the grammar. Scope changes and declarations that modify the grammar also update the automaton.

In [HKR89] a simple algorithm for incrementally updating the LR(0) automaton is given. Suppose that we are using the graph representation of the viable-prefix recognizing DFA. The idea is that each state is represented by its set of kernel items (see [ASU86]), the set of reductions, the sets of possible goto and shift transitions, and a “dirty” flag. Initially, only the start state exists, and it is “dirty”. When the parser executes and wants to examine the transitions from a particular state, it first checks the dirty-flag and if set, computes the closure of the state. This in turn may cause new initial “dirty” states to be created, or links to be made to existing states.

The incrementality here means that changes to the grammar only need to do minor adjustments to the cached automaton. It is not until syntactic objects are being used that the real generation work begins. Once completed, the automaton is reused until further changes to the grammar causes new incremental updates.

4.4.4 Pseudo-parallel LR(0) parsing

It is generally believed that LR(1) or some of its approximations is needed to build deterministic parsers for standard programming languages. Unfortunately, LR(1) automata are considerably more expensive to build than LR(0) automata. A technique known as pseudo-parallel LR parsing was first developed for natural language processing by Tomita [Tomita85]. It achieves the recognizing power of LR(1) by running several LR(0) parsers in parallel. The idea is this: suppose that an LR(0) automaton in a certain state finds
some shift/reduce or reduce/reduce conflicts. Instead of giving up, it can be cloned, and each copy can be set off to pursue one of the alternatives. In practise, a single LR-engine maintains a set of parsers synchronized by all being at the same position in the input.

This approach has two advantages. First, the overhead of pseudo-parallel parsing is low enough to be practically useful when short sentences are involved (within a factor of two compared with LALR(1) on sentences with up to several hundred tokens [Rekers92]). Secondly, the method automatically handles the case when ambiguous grammars are involved, and the system wants to find all the parse trees rather than a single unique one.

The DML system uses both the incremental and pseudo-parallel methods. The latter fits nicely with DML’s way of first generating all parse trees, and then removing the type-inconsistent ones. Currently the algorithm is based on [HKR89] which limits grammars to the “finitely ambiguous” context-free grammars (there must be finitely many parse trees for any given sentence).

4.4.5 LR(0) versus SLR(1) and LALR(1)

Why LR(0)? Would not SLR(1) or LALR(1) be more efficient? Not necessarily. The advantage of LR(0) is that no lookahead information needs to be computed. The disadvantage is that lookahead could be used to prune the number of generated parsers, instead of blindly trying all possibilities (in the case of shift/reduce or reduce/reduce conflicts). However, in [Lankhorst91] it is reported that SLR(1) and LALR(1) only buy about 10% in parsing time over LR(0). Note also that lookahead would have to be recomputed in some cases, as in the example below:

```plaintext
syntax t1 = "foo"
and t2 = "(" t1 ")" (* FOLLOW(t1) = {"\n"} *)
let
  syntax t3 = "x" t1 "x" | "z" (* FOLLOW(t1) = {"\n"}, "x" *)
in
  ...
end (* now, FOLLOW(t1) = {"\n"} again *)
```

5. Interaction with the ML module system

Syntax declarations have an unfortunate interaction with the module system. Ordinary datatype constructors can be accessed either by opening a module, or by using the long identifier syntax <module>.<name>. Since there is no way to refer to the constructors of a syntax declaration except by using syntactic objects, one must open the module in order to bring the constructors and the grammar environment in scope. An alternative would be to allow the user to explicitly name the constructors, for example:

```plaintext
syntax t = "foo" : Foo
   | "bar" : Bar
```
This alternative solution has not been investigated yet.

6. Conclusions

To our knowledge this is the first both general and efficient approach to concrete syntax objects within ML. An implementation, as presented in this paper, has been done within the DML system. The primary applications so far has been in denotational language specifications, including a C subset and full Scheme, from which compilers and interpreters have been generated. Traditional compilers not based on formal semantics have also been implemented. In all of these cases, the use of syntactic objects (mainly patterns) for instead of conventional data types was found to make the specifications easier to read and write. Some care must be taken with highly ambiguous grammars; in these cases some strategically placed explicit type constraints always suffice to make the resolution successful. The implementation, although being a prototype, has been fast enough not to present any problems.

7. References


An optimizing ML to C compiler

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Abstract

Since the C language is a machine independent low-level language, it is well-suited as a portable target language for the implementation of higher-order programming languages. This paper presents an efficient C code generator for Caml-Light, a variant of CAML designed at INRIA. Fundamentally, the compilation technique consists of translating ML code via an intermediate language named Sqil and the runtime system relies on a new conservative garbage collector. This scheme produces at the same time excellent performance and good portability.

1 Introduction

ML is a complicated programming language. Its syntax is complex and its dynamic semantics is related to the semantics of lexical Lisps like Scheme (cf [IEE90]). Furthermore it offers a strong polymorphic type system and a safe module system. Because of its overall qualities this language has become more and more popular for research and teaching, but with the emergence of better compilers it could reach a larger audience in industry as well.

At this time there are a few good implementations of ML, but all suffer at least from one or another drawback: disappointing execution speed, lack of portability or excessive use of memory.

The existence of a static type system in ML should permit us to compile this language nearly as efficiently as Pascal (not quite because polymorphism requires all objects to be the same size). The choice of C [KR88] as target language allows us to get a portable compiler easily because C is so widespread and now well standardized; nevertheless this choice must not cause bad general performance. Using the K2 compilation kit of Nitsan Séniak and Vincent Delacour, we built a very efficient and machine-independent ML to C compiler [Cri91].

In this article we explain our approach to ML compilation, provide a sketch of our implementation and give some results. We use the word 'global' to mean imported or exported and its contrary 'local' to mean local within a given module or function.

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2 Compiling strategy

We use a new compilation technique, named *explicit specialization* [Sén89], to obtain very good optimization of function calls: the cost of function calls is indeed the bottleneck inherent to the compilation of all functional languages.

Our compiler reduces the semantic complexity of ML programs, according to some simple and well-defined steps, to a level acceptable by a compiler with a familiar technology like a typical C one. As a matter of fact people are accustomed to writing programs that use mostly the capabilities of classical algorithmic languages such as Pascal. In this case, the dynamic usage of local functions is limited to the static scope of their definitions, which fortunately allows us to handle functional values without allocating a closure in the heap: well-known and efficient techniques (cf [ASU86]) can be used to compile classical algorithmic languages.

Since data is handled in the obvious manner in our translation scheme and is later optimized sufficiently by modern C compilers, we concentrate our efforts and description on control flow.

2.1 The intermediate language: Sqil

The specialization technique consists of translating ML programs that use general control flow constructs into more specialized and better compilable forms, the Sqil ones.

The Sqil dialect, kernel of the K2 compilation kit, is a kind of lexical and bivalued¹ Lisp with continuations and functional values restricted to the global environment. Since Sqil only addresses the efficient compilation of control flow, it is limited to a few elementary special forms:

- `defun`, `defvar`, `progn`, `let`, `if`, `setq`, `labels` and `flet` are present with the same meaning as in Common Lisp (cf [Ste84]);
- `(function symbol)` yields the functional value of `symbol`, which must be a global function. This value can be later used in a computed call with `funcall`;
- `(the-continuation)` yields the calling continuation of the global current function definition. This continuation can be later invoked by `(continue cont value)`;
- finally, `(block name expressions...)` and `(return-from name value)` build lexical escapes.

The restriction on functional values and continuations guarantees one important property: identity between variables and registers (see [Sén91]), meaning that each local variable of a program can be allocated to a register. Additionally function call is direct by default thanks to bivaluation; computed calls explicitly use `funcall`.

We successfully used the K2 compiler written by Nitsan Séniax, who has developed original techniques to compile local functions towards C [Sén90]:

- *Displacement of continuations* propagates calling continuations of functions into the called functions in order to transform many non-tail calls into tail ones, which can be compiled using `goto`.

¹I.e. symbols can have a functional value and a non-functional one at the same time, like in Common Lisp.
• Function integration in another function definition allows K2 to compile mutually
tail calls between several functions into ordinary jumps.

The K2 compiler has proved to yield very good C code. Nevertheless Sqil could be directly
compiled to efficient native code as well.

2.2 C as a target language

The Common Lisp compiler AKCL [Sch88] shows that it is realistic to compile Lisp or
ML towards C used as a portable target language. Actually the primary advantages of C are:

• C is a low-level language, whose compilation captures all previously performed high-
level optimizations well.

• C is highly portable and available on an increasing number of architectures, partly
thanks to the spread of the Unix system.

• Thirdly, C compilers progress each year, while good assembly code generation by
hand for RISC architectures is becoming very complicated.

However, in some respects, C is not so well-suited as a target language. Here are some
reasons:

• The compiler designer loses full knowledge and mastery about the generated ma-
chine code: for instance, some critical optimizations concerning the function calling
sequence that are performed by some Lisp compilers cannot be carried out any more.
Moreover multiple results can be inefficiently compiled.

• Some specialized machine code instructions cannot be generated by the C compiler,
like arithmetic operations with overflow checks. But this is less relevant for RISC
architectures.

• Moreover, the arrangement of local variables in the C stack or in registers is left
undefined. We cannot therefore use a classical garbage collector because it requires
us to identify precisely all roots that can point to allocated data. Consequently we
shall use a more complicated conservative garbage collector with ambiguous roots.

3 Implementation

3.1 The front-end

Our front-end is much the same as ZINC's one, written by Xavier Leroy [Ler90]. It first
parses the Caml-Light grammar and then infers the types of the expressions using a variant
of Milner's algorithm [Mil78]. The typechecker stores type information about each symbol
in the global environment, which enables us to use the type of global functions during the
code generation pass.
3.2 Code generation

3.2.1 Pattern-matching compilation

The pattern-matching compilation pass translates ML pattern-matching into a tree of Sqil's case selection instructions that is later translated to switch in C. The inherent backtracking involves the use of Sqil's lexical escape blocks, which generally compile to goto. This translation exactly follows the algorithm given in [Pey87], which is easy to implement but not optimal.

3.2.2 From ML abstract syntax tree to Sqil

We need two local environments for each toplevel phrase in this translation step:

1. An environment containing each variable with its access path from root to node in its defining pattern.

2. A local function environment containing the arity of each function name.

According to the different top nodes of the abstract syntax tree the translation proceeds in various ways:

• For the case of an identifier, we must distinguish a local or a global identifier, referring to a variable or to a function, because Sqil discriminates both kinds of values.

• For the case of a type constructor, we try to propagate constants, whenever it is possible thanks to the immutability of ML data types. Unfortunately this cannot be done with vectors and character strings that are inherently mutable in Caml-Light.

• The application case is the most complicated, because we want to uncurry most function calls:
  
  - When applying a local function or a primitive to a number of arguments matching its arity\(^2\), we translate it to a direct application. But if some arguments are missing, we must add the code to build a partial function (using Sqil's function form); and if there are too many arguments, we have to compute additional calls using funcall.

  - The handling of global functions differs because the actual arity of an imported function is not accessible: only its type is known, and gives us its maximal arity. In order to increase execution speed, we insert several entry points for each global function, ranging from one argument to the maximum arity deducible from the function type. Thus a global function application simply calls the entry point corresponding to the number of available arguments.

  - If the functional value of the applicand is to be computed at runtime, we iterate funcall on each argument, since all functional values, i.e. closures, are of arity one in our scheme.

• The specialization of let needs the pattern-matching compilation pass and then distinguishes between variables, functions and recursive functions to be able to use let, flet and labels Sqil bivaluated forms.

\(^2\)We define the arity as the number of arguments available when the evaluation of the function can begin.
• The CAML try ... with exception block is compiled by storing in a global variable the corresponding continuation, after having saved the old one. This permits us to raise an exception dynamically just by invoking the last created continuation.

• The translation of other constructs such as conditional forms or sequential forms is straightforward.

Finally we must remember to translate global definitions of functions and variables into defun and defvar Sqil forms.

3.2.3 The globalization pass

Up to this point we have not paid much attention to the fact that continuations and function values of Sqil are restricted to the toplevel environment. That ensures that these are simple addresses of either a stack frame or a global function entry.

If this is not the case, we have to globalize the function: a local function \( f \) that is referenced as a functional value must be moved to the toplevel and defined there with an additional argument representing its environment, i.e. its free variables. Then in general it is necessary to build a closure each time we need the functional value of \( f \), and send its current environment together with its argument whenever \( f \) is called.

Moreover each free function of \( f \) will be either integrated in \( f \) if it is only referenced in \( f \), or else must be globalized too. So any free variables of these functions must be present in \( f \)'s environment too.

If a local function contains the form (the-continuation), it must be globalized as well, and additional arguments must be supplied for its free variables.

3.2.4 C code expansion and linking

When compiling Sqil code to C, remember that such optimizations as elimination of tail-recursive calls, continuation displacement and integration of local functions are performed. Macros for data handling are expanded too.

The Caml-Light module system is simple but quite natural and is based on interface files that provide the names and types of all exported global identifiers of a module. It allows separate compilation between modules and the definition of abstract types such as polymorphic hashtables. So we have to link all compiled modules (along with the runtime support) to obtain the executable file.

The standard library is entirely written in ML; it provides C written primitives only for low-level tasks such as I/O or Unix system calls.

3.3 The runtime system

3.3.1 The selected memory model

All ML objects are uniformly represented by a machine word. Characters, integers, short reals and some special sum types such as booleans are immediate values whereas all other objects are statically or dynamically allocated in several words and represented by a pointer. We discuss below whether pointers and immediate values need to be distinguishable.

Closures and environments are simply allocated as a vector in consecutive memory words. So extending an environment by new variables is not implemented by chaining,
but needs to copy the values into a bigger environment vector. Sum types require a small integer tag for the purpose of discriminating different constructors.

3.3.2 Which is the best garbage collector?

In order to collect memory which is no longer used, we need a garbage collector that can trace ambiguous roots in the C execution stack. It is possible to use a mark and sweep algorithm in a BIBOP memory model without any tags in the objects themselves. With this scheme all fields in an allocated object are ambiguous.

We have tried this scheme and used Boehm’s portable collector [BW88] for a while, but the results were not very good for programs that allocate a great amount of data in the stack or in the heap. The garbage collector was indeed spending too much time in its mark function because it considers every word in the stack or in the heap as a potential pointer.

We think it is much better to distinguish pointers and immediate values with a tag. First the difference between, say, an integer and a pointer in the heap and even in the stack is much more immediate. Then one can use a mostly copying algorithm, such as Delacour’s one [Del91], which is more efficient because it does not scan the whole heap like a mark and sweep algorithm and because it increases locality by copying and compacting the structured objects.

Moreover the runtime system can easily offer the user ‘generic’ functions such as equal that need to explore the tree structure of the objects, and give these functions a polymorphic type.

Our implementation uses a one-bit tag scheme that renders all immediate values even and all pointers odd. This tag scheme gives better results than the opposite one, since arithmetic operations are little altered while pointer dereference only needs a small offset, which is free on many machines.

3.4 Benchmarks

<table>
<thead>
<tr>
<th>Compiler</th>
<th>SML-NJ</th>
<th>Caml-Light</th>
<th>CeML</th>
<th>Zinc K2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Version</td>
<td>0.75</td>
<td>0.41</td>
<td>without GC</td>
<td>with GC</td>
</tr>
<tr>
<td>Sieve</td>
<td>4.2</td>
<td>11.6</td>
<td>2.4</td>
<td>1.1</td>
</tr>
<tr>
<td>Solitaire</td>
<td>21.3</td>
<td>124.6</td>
<td>7.4</td>
<td>4.7</td>
</tr>
<tr>
<td>Takeushi</td>
<td>18.7</td>
<td>49.6</td>
<td>3.5</td>
<td>6.1</td>
</tr>
<tr>
<td>Tak w. exceptions</td>
<td>33.0</td>
<td>73.6</td>
<td>77</td>
<td>45.2</td>
</tr>
<tr>
<td>Knuth-Bendix</td>
<td>1.9</td>
<td>8.8</td>
<td>12</td>
<td>2.7</td>
</tr>
<tr>
<td>Boyer</td>
<td>5.6</td>
<td>22.7</td>
<td>n.a.</td>
<td>3.9</td>
</tr>
<tr>
<td>Euclidian division</td>
<td>3.4</td>
<td>20.2</td>
<td>17.5</td>
<td>2.7</td>
</tr>
<tr>
<td>Church integers</td>
<td>5.5</td>
<td>31.3</td>
<td>25.2</td>
<td>3.8</td>
</tr>
<tr>
<td>List summation</td>
<td>24.8</td>
<td>41.4</td>
<td>10.6</td>
<td>6.1</td>
</tr>
<tr>
<td>Integral</td>
<td>34.1</td>
<td>59.7</td>
<td>14.2</td>
<td>5.5</td>
</tr>
</tbody>
</table>

Figure 1: Comparison among some ML compilers (user times in seconds)

Some benchmarks are listed on figure 1. They were performed on a Decstation 3100 with the R2000 RISC processor. Tests on a Sparc architecture yield similar results.
We have chosen three other good compilers to make some comparisons. These are the well-known SML-NJ native code compiler [AM87], the fast bytecode interpreter Caml-Light and another ML to C compiler, Emmanuel Chailloux's CeML [Cha91]. Roughly speaking, SML-NJ uses a Continuation Passing Style translation scheme with a fast stop and copy generational garbage collector, and allocates everything on the heap [App87]. Caml-Light uses a similar runtime as ours, but its collector is a non-conservative one with generations, whereas CeML has a good conservative mark and sweep algorithm, but scans a special-purpose application stack instead of the C stack and uses a type tag in the objects themselves.

The first tests - Erathostenes' Sieve, the game of Solitaire and Takeushi's function - show that the C generated code is at its best when compiling imperative programs. That was indeed expected! The next two - Tak using exceptions to return each partial result and the Knuth-Bendix completion algorithm - demonstrate that exception handling is quite expensive in C, because you generally have to use the library functions setjmp and longjmp.

Knuth-Bendix completion procedure, Boyer's tautology checker, Euclidian division (extracted from the Coq proof assistant) and Church integers are very functional programs. Zinc-K2's excellent performances with respect to these programs show that with our optimizing compilation strategy C code generation can achieve the best performance results for typical ML programs.

List summation tests list processing whereas Integral tests floating point arithmetic. Except for this last test, garbage collection is cheap (actually the version without a GC uses a simple allocation fringe pointer and objects have no tag). The reason for this overhead is that we cannot tag short reals, so we have to box them. An alternative would be to adopt a more clever boxing mechanism (see [Ler92]).

4 Conclusions

We have demonstrated that C code generation can be very efficient if one succeeds in making a good use of the control flow instructions present in C, despite its bad exception handling mechanism. Uncurrying is a critical factor in achieving this, because it eliminates a lot of redundant partial applications. The use of an appropriate intermediate language permits us to describe the relevant optimizations easily.

The next step would consist of giving an object its natural C type whenever it is known statically to have a monomorphic ML type. This would allow the C compiler to handle the object directly, yielding a better assembly code, and would avoid many useless heap allocations. We plan to develop this idea and to add pertinent datatypes to Sqil.

Finally, a garbage collector with ambiguous roots can be (nearly) as efficient as a conventional one, and a clever runtime tag mechanism does not hamper the overall performance.

5 Acknowledgements and related works

We would like to thank Xavier Leroy for his precious help, Emmanuel Chailloux for some valuable discussions, and Bruno Monsuez and Alan Mycroft respectively for their remarks about the first and second draft of this paper.
Previous work on the compilation of Lisp to C include the Kyoto Common Lisp [YH88] and Bartlett's SCHEME→C compiler [Bar89]. A previous attempt to compile ML to C is reported in [TAL90], but SML2C yields lower-level C programs, for example it does use an apply-like procedure for function calls instead of the standard C calling mechanism.

References


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An Efficient Way of Compiling ML to C

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Introduction

The ever increasing diffusion of ML has yet to go beyond the areas of education and research. To change this situation, two conditions must be satisfied: in their shared characteristics, ML programs need to become as efficient as imperative programs without sacrificing security for efficiency and must also be easily interfaceable with other libraries. I propose a new compilation model for ML, called CeML [6], derived from the CAML [17] (ML dialect), which translates ML directly into C (understood as a portable assembler). In fact an abstract machine, inspired from FAM [15], is partially included in a runtime library. A particular attention has been paid to two points. The first one is function application which is optimized using an extended notion of functional type. The second one is the correspondence between basic ML types and basic C types. This allows ML objects including structured types, represented by pointers, to directly use the call protocol of C. This correspondence creates some difficulties for the Garbage Collector, but they vanish thanks to a new Mark&Sweep [12] with ambiguous roots [4], where basic values are not tagged.

All these implementation features are independent, making no supposition regarding the C compiler because they use another stack for the root set and for the application. In this manner, C programs created by CeML are easily interfaceable.

1 CeML Definition

CeML is a ML dialect derived from CAML language [17]. It differs from CAML with regard to its type system (with the introduction of $\Rightarrow$, a new functional type constructor) and its module system.

The main syntactic restriction is the absence of local declaration for type and exception.

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1.1 Type system

The λ-calculus has no notion of abstraction arity. The following notation: \( \lambda x y . x y \) is just a convenient way of writing \( \lambda x \lambda y . x y \). But in a programming language we need to have this notion in order to obtain an efficient application. In CeML, we can define the function arity, bound to a function name, if there is no intermediate evaluation or local declaration between two (\( \lambda \)) (function). For example:

```plaintext
let addition = function x -> function y -> x+y;;
```

becomes a function with two arguments, and its type is \( \text{int} \rightarrow \text{int} \rightarrow \text{int} \). This type is different from \( \text{int} \rightarrow \text{int} \rightarrow \text{int} \rightarrow \text{int} \rightarrow \text{int} \rightarrow \text{int} \), which represents the classical functional type constructor. To formalize this type system, we give the following typing rules:

(App)

\[
\frac{C \vdash \text{expr}_1 : \tau' \rightarrow \tau \quad C \vdash \text{expr}_2 : \tau'}{C \vdash \text{expr}_1 \text{ expr}_2 : \tau}
\]

(TApp)

\[
\frac{C \vdash \text{expr} : \tau_1 \rightarrow \ldots \rightarrow \tau_n \rightarrow \tau \quad C \vdash \text{expr}_1 : \tau_i (1 \leq i \leq n)}{C \vdash \text{expr} \text{ expr}_1 \ldots \text{expr}_n : \tau}
\]

(PApp)

\[
\frac{C \vdash \text{expr} : \tau_1 \rightarrow \ldots \rightarrow \tau_n \rightarrow \tau \quad C \vdash \text{expr}_1 : \tau_i (1 \leq i \leq k \text{ with } k < n)}{C \vdash \text{expr} \text{ expr}_1 \ldots \text{expr}_k \rightarrow \ldots \rightarrow \tau_n \rightarrow \tau}
\]

(E)

\[
\frac{C \vdash \tau' \rightarrow \tau}{C \vdash \tau' \rightarrow \tau}
\]

(I)

\[
\frac{C \vdash \text{pat}_i : \tau_i \quad \text{expr} : \tau}{C \vdash \text{function pat}_1 \rightarrow \ldots \rightarrow \text{function pat}_n \rightarrow \text{expr} : \tau_1 \rightarrow \tau_2 \rightarrow \ldots \rightarrow \tau_n \rightarrow \tau}
\]

The great advantage of using a new type constructor to detect curried functions is the type propagation of the function result. The following function:

```plaintext
let add4 x y = let f u v = x+y+u+v in f;;
```

has this type: \( \text{int} \rightarrow \text{int} \rightarrow (\text{int} \rightarrow \text{int} \rightarrow \text{int}) \) which indicates the functional arity of the result.
remark In ML, user defined polymorphic function can explore the structure of a polymorphic parameter. In CeML, this rule is also applied to primitives. This is the reason why the symbol = has the semantic of eq (equality for immediate values or sharing for structured objects). It is a problem for the programmer but it is more coherent for the language definition because the ML polymorphism is parametric i.e. it does not look at the form of the function arguments.

For example, CeML includes a parametric polymorphism, = thus means eq i.e. describes an equality for immediate values or a sharing for structured values. Effectively = cannot mean equal because with parametric polymorphism a function does not explore the structure of a polymorphic parameter [14].

1.2 Module System

The CeML module system is not so powerful as CAML or SML module systems. It is only a convenient way to make separate compilation. Separate compilation is a convenient way of dividing programs into different compilation units. A CeML unit is decomposed into three parts: importation, implementation and exportation. The importation part describes the sub-graph of dependences at level one. The complete graph for a program has no cycles. The gray arrow of figure 1 is forbidden.

![Dependencies graph](image)

Figure 1: Dependencies graph

2 C as Intermediate Language

Many compilers consider C as a portable assembler. For example: some imperative languages (f2c, p2c), object languages (C++, Objective C, Eiffel, Modula3), logical languages (WAMCC) or functional languages (Scheme->C, KCL, SML2C, K2C, Z2K2C). These language references can be found in [6]. I present the main advantages and inconveniences to use the C language as assembler.

2.1 Advantages and inconveniences

2.1.1 advantages

- intrinsic: The memory allocation is sufficiently low level that it permits many kinds of manipulations. Global functions authorize many arguments, and control structures are powerful (loops, jump function for switch).

- standard: There is a standard ANSI C. The other C syntax (K&R [10]) is similar.
efficiency: A great effort is effectuated by hardware manufacturers to obtain excellent C compilers on new processor architectures (RISC). One of the most popular benchmarks is the SPECMARK that gives a coefficient of a C compiler for a processor.

tools: Many tools permit the profiling and debugging of the generated C code.

2.1.2 inconveniences

The main inconvenience in using C as an intermediate language is the absence of a good exception system. For example, C neither detects integer overflow nor C stack overflow.

2.2 What must be added to C to compile ML?

We need the following extensions to compile CeML to C:

- a Garbage Collector
- a general application mechanism
- an exception handling

These different extensions will be introduced into a runtime library that will need to be linked with C generated programs.

3 Compilation Scheme

The compilation scheme is classical, but its presentation gives a global vision.

For each CeML compilation unit (in fact a file with .ml extension), there is a corresponding C file (figure 2). After this compilation phase, the link phase (figure 3) creates an executable program from different units and from the runtime library.
CeML is actually a CAML program. For the computers supporting CAML it can be used directly as a command line. For the others, the C files must be generated on another computer and then be compiled on the host computer.

4 Runtime Library

In terms of efficiency, the runtime library is one of the essential parts of the implementation. I shall describe the type representation and a new Garbage Collector for the memory management, the application mechanism and the exception handling.

4.1 memory management

In ML, with a static typechecker, tags are effectively unnecessary [1]. In fact, only distinguishing information is needed for the summation types in order to distinguish constructors of a same type and the different kinds of vector. We follow this rule so as to be close to C basic types.

4.1.1 data type representation

Immediate values and pointers have a 32 bits representation. Others values (accessed via pointers) are represented in the heap by various structures. In order to distinguish between them, some type information is required.

Integers and floating point numbers use a word as well. Double precision floating point numbers are not implemented, but they can be represented as a pointer toward a four word storage.

4.1.2 Garbage Collector with ambiguous roots

I present a new Mark&Sweep algorithm with ambiguous roots. I describe the data representation, the partitioning of memory, the setroot and the algorithm used during the Mark phase to distinguish immediate values and pointers.

partitioning memory The heap is partitioned into chunks (cf. figure 5). Each chunk contains objects of the same size (to the power of two). There are \( nbzones \) sets of chunks called zone (from \( 2^1 \) to \( 2^{2+nbzones} \) bytes). Objects greater than one chunk are arranged into several chunks. This partitioning of memory is a variant of the BIBOP
(Big Bag Of Pages [15]) algorithm. For our implementation, the chunk size is four kilobytes and $nbzones$ is equal to ten.

**Figure 4: Data type representation**

**Figure 5: Partitioning memory**

**free lists** Each predefined zone has a list of available elements (cf. figure 6).

**Figure 6: Free Lists**
setroot  The setroot is represented by a statically allocated independent stack. This stack allows for the storage of immediate values or pointers. Sometimes there is a double use between this stack and the C stack, but this choice is necessary so that the former stack may be independent of the C stack. This stack is also used for the general apply mechanism (when a direct call to a C function is not possible because the argument is a closure, or during a partial application).

initial memory state  In the beginning, the heap is empty. Each zone can grow dynamically. The first allocation for a zone is ten chunks. The next allocations are computed by the growing function, after the Sweep phase. This orientation permits to control the heap evolution.

allocation  There are two kinds of object allocation. The first one is used for small objects less than one chunk in size. This first case has two alternatives. When the object size is known (for example one cons uses four words) then, if the corresponding free list is not empty, the allocation is completed, otherwise a GC is invoked. In other cases the zone to be used has to be computed. If the object size is greater than a chunk, then the object uses several contiguous chunks.

recovery  When a zone is full, one must recover some space. There are two phases, the first (Mark) marks each object indicated by the setroot and the second (Sweep) preserves only these objects.

mark  For each value inside the setroot, a discriminating algorithm distinguishes between an immediate value and a pointer. In this case the structured object is marked and the process is applied to its structure elements. This algorithm is recursive, but it does not use the heap. Instead the recursive calls are pushed into the C stack.

sweep  For each chunk in use, its corresponding free list is updated by all the unmarked elements. This algorithm explores all used memory. This is an implementation which wants to be simple. If the responsible zone which raised the GC is too small after memory recovery, then new chunks are allocated for this zone.

distinguishing algorithm  When the GC examines a memory cell, it decides that its content should be considered as a pointer and marks the pointed object if the four following tests succeed:

- is the pointed object in the heap?
- does this address belong to a chunk in use?
- is the pointed object correctly aligned for this chunk?
- is there an object allocated to this address?

Otherwise, the content of the memory cell can be safely considered as an immediate value.
It is important to use a GC which displaces no objects, for the correspondence between CeML variables and C variables because otherwise the variable storage into the CeML stack is more costly (in this latter case two kinds of storage, one for C values and another for C variables are needed, and the pattern variables must also be stored in the root set).

4.2 General application mechanism

The general application mechanism uses the CeML stack, in order to push a closure and arguments. A closure is represented by a pair: (code,environment) where the code is a C function pointer and the environment a vector. More data is necessary in the closure, for example the maximum number of arguments (including environment values). The application expects a closure and i arguments on the top of the stack (cf. figure 7).

![Figure 7: Application using stack](image)

The return value of this application is considered as a C value, and returns to this evaluation step.

4.2.1 total application

The body evaluation of a function occurs when all its arguments have been passed. We note $C^n_p$ a closure with $n$ variables where $p$ are already given. The total application begins if there are $i = (n - p)$ new arguments on the top of the stack. Let $fn$ be the C function associated in the closure, the return value is:

$$fn(env[1], env[2], \ldots, env[p], Stack[SP+1], Stack[SP+2], \ldots, Stack[SP+i])$$

There is no closure duplication if all arguments are given.

4.2.2 partial application

In this case the environment portion of the closure is duplicated. The environment vector increases as in figure 8. For each partial application there is a closure duplication.
4.3 Exceptions

CeML exceptions are considered as constructors, but this type is open.

The runtime library uses the C library setjmp and longjmp. setjmp is used for the try and longjmp for the raise. But it is necessary to add data to the C context (jmp_buff,) such as the CeML stack pointer. The return value from a raise is stored inside the stack. This stack pointer must be preserved in the CeML context. All the different CeML contexts are inside a linked list (EP is the head), as below:

5 Code Generator

I shall explain only the main features of declaration and expression generation. There is a direct correspondence between CeML variables and C variables. During the generation phase, the CeML compiler uses four contexts, described below:

- stack context : is true if the expression must be pushed.
- variable context : has a variable name if the expression must modify this variable.
- return context : is true if the expression must be returned by a C function
- application context : is a list of composed applications.
I note that for $[expr, return, variable, apply]_{stack}$ or $[expr, return, variable, apply]_{env, arity}$ for the compilation of expressions in a compilation context. For the declarations, I write $[decdvar]_{env, arity}$ for a declaration with env as environment and arity as function arity. For example, a non functional variable has a zero arity, and global variables always have an empty environment. In all cases, a C eML variable is translated by one or two C variables.

5.1 non functional declarations

C authorizes local variables except for the functions, but closures have a C type defined as Closure. Then the translation for CeML variables of any type, except $\Rightarrow$, is direct.

$$\{let \ v = e\}_0 = \{typeC \ v ; \ \{e\}_{true, false, \ "v" = \ "\"} \}$$

where typeC is the corresponding C type for the CeML type.

For the local and non functional declarations, the translation is identical because the local environment is known at this level.

5.2 Functional declarations

5.2.1 global declarations

In this case, the environment is empty. Then the number of parameters of the corresponding C function is the same as in CeML. In the following example, the $f$ function has the type $\alpha \Rightarrow \beta \Rightarrow \alpha$.

$$\{let \ f = e\}_{0, 2} = Closure \ _M.f ;$$

$$Ptr \ _F.f(\_V.1, \_V.2)$$
$$Ptr \ _V.1 ;$$
$$Ptr \ _V.2 ;$$
$$push\_safe(\_V.1) ;$$
$$push(\_V.2) ;$$
$$\{ \ _P.r \ ; \ [e]_{false, false, "R" = \ "\"} \}$$
$$pop\_n(2) ;$$
$$return(\_P.r) ; \}$$

There are two C variables for each CeML function. One for the closure data structure and a second for the C function.

The previous example also initializes a C closure ($\_M.f$) with its corresponding C function pointer ($\_F.f$) and its arity (2), as follows:

$$\{let \ f = e\}_{0, 2} = \_M.f = init\_closure("filename", "f", \_M.f, \_F.f, 2) ;$$
$$push(\_M.f) ;$$
5.2.2 local declaration

The binding of local function free variables is resolved by adding extra parameters to the C function, using $\lambda$-lifting [9]. Each free variable creates a new parameter, but it is not enough because a free functional variable can contain others free variables. The free variable set contains all variables not bound under a $\lambda$ or a pattern-matching, and not defined at the global level. It is necessary to build a dependence variable set ($DV$) from the free variable set ($FV$) as follow:

$$DV_0(v_i) = FV(v_i)$$

$$(k \geq 0) \text{ if } DV_k(v_i) = w_j \text{ then } DV_{k+1}(v_i) = \bigcup DV_k(F\{w_j\}) \cup V\{w_j\}$$

where $F\{w_j\}$ and $V\{w_j\}$ represent respectively the functional variables and the non-functional variables. of $w_j$, then $DV = DV_{k+1}$ if $DV_{k+1} = DV_k$.

The great difference between non recursive and recursive declarations comes from the initial free variable set.

**non recursive declaration** For the following CeML sentence (E) : let $v_1 = e_1$ and $v_2 = e_2$ ... and $v_n = e_n$ in $e$ we obtain:

$$FV(v_i) = FV(e_i)$$

$$FV(E) = \bigcup_{i=1}^{n} FV(v_i) \cup (FV(e) - \{v_i\})$$

**recursive declaration** For the following CeML sentence (E) : let rec $v_1 = e_1$ and $v_2 = e_2$ ... and $v_n = e_n$ in $e$ we obtain:

$$FV(v_i) = FV(e_i) - v_i$$

$$FV(E) = \bigcup_{i=1}^{n} FV(e_i) \cup FV(e) - \{v_i\}$$

**translation** Then the translation uses the dependence variable set :

$$[\text{let rec } v_1 = e_1 \text{ and } v_2 = e_2 \text{ in } e]_{s,r,v,l} = [\text{let rec } v_1 = e_1]_{(DV(v_1),n1)}$$

$$[\text{let rec } v_2 = e_2]_{(DV(v_2),n2)}$$

$$[e]_{s,r,v,l}$$

All the variables belonging to $DV(v_1)$ are translated as extra parameters to the corresponding C function.

5.3 Application optimization

The general case application is the following:

$$[expr_1 \text{ expr}_2]_{s,r,v,l} = [expr_1]_{true, false, \cdots} [expr_2]_{true, false, \cdots} apply(2)_{s,r,v,l}$$

where $true$ in stack position indicates that the expression must be pushed.
5.3.1 total application detection

The idea is to compare the function arity and the number of arguments given. If \( p \) arguments are given to \( f \) then:

- if \( f \) type is \( t_1 \rightarrow t_2 \) then it is the general case.
- if \( f \) type is \( t_1 \Rightarrow t_2 \Rightarrow \ldots \Rightarrow t_{n+1} \) then there are two sub-cases:
  - \( p < n \) then \( f \) followed by the \( p \) arguments are pushed and \( \text{apply}(p+1) \) is called.
  - \( p \geq n \) then \( f \) is directly called with the first \( n \) arguments. If some arguments remain then another application is efectuated.

5.3.2 optimized application

Here we witness the use of C protocol call functions.

\[
[f \ e_1 \ldots \ e_n]_{(s,r,v,l)} =_{\text{opt}} \{ \text{typeC} \ t_1 \ T_1; \\
\ldots \\
typeC \ t_n \ T_n; \\
[e_1]_{\text{false,false},"T_1"} \text{push}(T_1); \\
\ldots \\
[e_{n-1}]_{\text{false,false},"T_{n-1}"} \text{push}(T_{n-1}); \\
[e_n]_{T_n} = \text{pop} \ n(n-1); \\
F\_f(T_1,T_2,\ldots,T_n)(s,r,l,v) \}
\]

5.4 Pattern matching

For all types except integers and summation types, a naive algorithm (sequence of if) is used. For the integers and the summation types a switch control structure is used after a reorganization of the pattern matching. Constructors are represented by an integer. This optimization cannot be used with exceptions which belong to an open type.

6 Performances

I compare different ML compilers, divided into two families: CAML [17] and SML [13]. CAML and SML have a very similar core language. We can consider the following compilers compile the same core language. The CAML compiler translates to the CAM [7] code which is expanding into native code. CAML-LIGHT [11] translates to a byte code which is interpreted. SML/NJ [3] uses a CPS [2] model and produces native code. CeML and SML2C [16] generate C programs, but SML2C is the back end of SML/NJ and produces low level C programs, in contrast to that CeML generates high level C code.
All these benchmarks have effectively been run on the same machine (DecStation 3100 with a processor MIPS R2000) under the same conditions. All compilers, except CAML, create executable programs. CAML-LIGHT creates a byte code object but its loading is rapid. The `exportFn` function of SML/NJ produces a large executable program (a core image of approximately two megabytes). CeML and SML2C generate C programs that are passed on to the C compiler. The respective size of executable programs are reasonable (under one hundred kilobytes for small CeML programs, and about four hundred kilobytes for SML2C).

The following benchmarks try to test the different parts of the ML language. Some of them are directly inspired from Gabriel’s work [8]. Fibonacci, Takeuchi, Integral and CountStr use basic types as integers, floats and strings. Takeuchi is the curried form of the famous Takeuchi function. Reverse, SigmaMap, ItList and Sieve test the construction and access to lists inside functional programs. Church int and DivEuclid are very functional programs. The former carries out calculations on Church integers. The latter is the Euclidean division extracted from the Coq system. TakExcept is always the Takeuchi function but written with exceptions. KB is the Knuth-Bendix term rewriting system applied to the group completion. It is very functional and uses exceptions. SigmaVect and Soli_let work on vectors. Soli_let is the resolution of the “solitaire game”.

In figure 10, all the numbers represent user time on a Unix operating system (given in seconds). Bold numbers indicate the best results.

<table>
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Figure 10: Experimental results

CeML’s excellent performance with respect to imperative programs was expected. But its results on the functional programs, when they are close to the Lisp style, justify a posteriori, its implementation choices. Its less satisfying times on very functional programs are not really a hindrance because in general, a program is scarcely so functional.
Nevertheless, its performances on exceptions are really a problem if their use becomes a programming style.

Conclusion

In this paper, I have attempted to demonstrate that functional languages can obtain the same efficiency as imperative languages for their shared characteristics without sacrificing security for efficiency which would be a bad deal. This was globally verified. In many cases, the C program generated by the CeML compiler, is very similar to the equivalent handwritten C program.

But at the same time, CeML had to be as efficient as the best ML implementations. CeML performances compares very favorably with the best ML implementations. There are however two types of programs where CeML performances are not highly satisfactory: the very functional programs (close to λ-calculus as in the case of Church integers) and programs which use too much exception handling (as Takeuchi with exceptions). In the first case, the partial application and the application of closures given as arguments forbid the application optimization.

In the second case, the exception handling mechanism is above all dependent on the operating system and does not yield good results on an Unix system. But since the partial application and the intensive use of exceptions are in fact scarce, they do not put into question the CeML implementation choices.

In fact, with its more informative typechecker (arity of functions, expressions decorated with their types), the CeML compiler yields excellent optimizations for the total application and the manipulation of basic values. But theses optimizations are not always possible. Typically when the application depends on a functional argument, then its arity is lost for the application optimization. The execution speed then varies according to the ratio of non-optimized application / optimized application. Its GC, with ambiguous roots, avoids the tagging of immediate values. The distinguishing algorithm, between an immediate value and a pointer, slows the Garbage Collector down, but the benefits achieved through the uniform representation of data is, in most cases, greater than the slowdown.

One could also complain that the CeML stack replicate a part of the C stack. But for two reasons I think it is better to use an independent stack than only the C stack. First, it is necessary to be independent from the C implementation. The C stack is not specified in the C language definition. For this reason, we do not make any supposition about the C stack. Secondly, the content of the C stack is unknown and can be large. It is particularly important for the GC which scans this stack to mark objects. CeML loses time in building this stack but it can save time if many objects are inside the C stack. Although the GC needs to allocate chunks in good alignment, it accepts holes inside the heap, and nothing forbids mixed allocation between CeML and other C libraries.

C programs generated by CeML are readable. This property allows to use C tools to profile these programs and facilitates the debugging of the compiler.

The reduction in the difference of execution times between C and ML, and the interface possibilities with others libraries open the door for a more widespread use of functional languages as everyday programming languages.
References


Abstract

Standard ML of New Jersey (SML.NJ) is a Standard ML compiler suitable for porting to different architectures. The compiler consists of a front end written in ML, a back end also written in ML, and a runtime system written in C and assembler. The front end is relatively independent of the target machine. This makes it easy to port the system to other architectures.

SML.NJ has up to now been reserved for workstations running UNIX. Code generators are available for a number of processors and the runtime system has been adapted to different UNIX versions.

The purpose of this paper is to report a new code generator for the Intel 80386 processor and a revised runtime system which runs under Windows. The code generator described in this paper is almost independent of the runtime system and consequently of the operating system, so it can be used in other architectures based on the 80386 processor, such as OS/2 and XENIX.

1 Introduction

Standard ML of New Jersey [4] is an implementation of the programming language Standard ML [6]. So far, Standard ML of New Jersey (SML.NJ) has solely been available on machines that run under the UNIX operation system. The purpose of this paper is to report a port of SML.NJ which runs on the Intel 80386 processor under Microsoft Windows 3.0, one of the most common combinations for PC's.

Our port of SML.NJ consists of two parts:

1. Standard ML modules which the SML.NJ code generator calls in order to produce 80386 machine code.

2. A new runtime system written in C and 80386 assembly language.

The first part is largely independent of Windows, i.e. it could be used for generating 80386 code that would run under other operating systems. The second part was necessitated by the fact that the runtime system has to run under Windows, which is very different from UNIX.

The reason we wrote parts of the runtime system in assembly language is that SML.NJ generates 32-bit code but no 32-bit C compiler is currently available under Windows.
When a 32-bit version of Windows becomes available, a new runtime system, which more closely resembles the original runtime system, can be written.

Few prerequisites are required to read this paper: familiarity with ML is necessary and some knowledge about the Intel 80386 assembler is useful.

The rest of this paper is organized as follows: in Section 2 we review those aspects of the 80386/Windows platform that are of particular interest in this context. In Section 3 we report the new runtime system and in Section 4 we describe the new code generator. Finally in Section 5 we show an example.

Many of the technical details that are not covered in this paper can be found as comments in the source code. References are made to the relevant files, which are located in the sml distribution directory.

2 The 386/Windows platform

In this section we discuss those aspects of the 80386 and Windows, that are important in the context of SML.NJ. Windows causes troubles in implementing the system on a PC. This is because Windows still is based on the old 8086 memory model where the memory is divided into 64k segments. As explained in Section 2.2, this fits badly with the assumptions which SML.NJ makes about storage usage; these assumptions are described in Section 2.1. In Section 2.3 we describe how one can fit the two together.

2.1 Assumptions that SML.NJ relies on

The compiler operates with two basic datatypes — integers and pointers — both 32-bit\(^1\). Thus it is desirable that the architecture supports 32-bit data and addresses. The way memory is allocated and the way the garbage collection is done [1, 2] assumes that the ML code and data are located in a single continuous memory block. Since compiled ML code accesses variables and functions in the runtime system through 32-bit pointers and vice versa, the runtime system should be in the same logical memory space as the ML code/data.

The compiled ML code occasionally makes calls to the operating system, of which some are critical and other are of a more peripheral nature. That is, some of the system calls are used by the compiler itself whereas others just are for use in user applications. The critical system calls must be supported by the operating system, or at least it must be possible to simulate a corresponding action.

Languages like ML make heavy use of memory, so the operating system should provide some kind of virtual memory.

In the next two sections we will see how the 80386 and Windows meet these requirements. Only a brief overview together with the choices made will be given here. For a more elaborate discussion of the problems with choosing a memory model and how to build the system, the reader is referred to [5].

\(^1\)actually only 31 because the least significant bit is used as a tag bit to distinguish between integers and pointers [2].
2.2 The 80386

The 80386 processor has 32-bit registers for manipulation of code and data. It is possible to address up to $2^{32}$ bytes (i.e. 4 Giga bytes), and the built-in paging mechanism makes virtual memory possible.

The memory organization is of interest to us. The 80386 operates with two kinds of segments:

* **USE16**: these are 16-bit segments. An address consists of a 16-bit segment address and a 16-bit offset. Code running in this type of segments by default uses 16-bit addresses and 16-bit data. Segment registers contain indices into a descriptor table that contains information about the type and location of the segment. With these segments the code and data has to be divided into 64K pieces, causing the well-known problems with managing large programs.

* **USE32**: these are 32-bit segments. An address consists of a 16-bit segment selector and a 32-bit offset. Code running in this type of segments by default uses 32-bit data and 32-bit addresses. This makes it possible to have a flat 32-bit memory model where the memory consists of a single contiguous block.

The same binary machine code will cause different actions in the two types of segments. When using USE16 segments, the 386 operates much like the predecessor 80286, only now 32-bit registers for manipulation of data are available. When using USE32 segments the processor acts like a real 32-bit processor.

The 80386 operates with separate code and data. It is not possible to write into a code segment nor to execute code from a data segment. But by letting a code segment register and a data segment register point to the same physical memory, self-modifying code becomes possible. This is known as *segment aliasing*.

The instruction set is comprehensive enough to implement the abstract machine that the front end generates code for. There are some problems in that the 386 has inherited many of the peculiarities from its predecessors. Some registers are dedicated to special purposes in some instructions and different addressing modes are available in different instructions. For details, see [5].

In short, the 80386 has what is needed to support the compiler. Unfortunately, Windows does not make all its capabilities available to the programmer.

2.3 Windows

Although Windows in enhanced mode exploits some of the processors 32-bit facilities, it continues to adhere the segmented memory model. It is not possible to implement a Windows application using an exclusively flat 32-bit memory model. Windows itself relies on 16-bit segments and a Windows application must contain at least one USE16 segment to interact with Windows.

Considerations about implementing the system using the USE16 memory model only, are given in [5]. Here I'll just state that it is not possible to get a satisfactory system within that model. Instead I'll explain how to run 32-bit programs under Windows.
The dynamic-link library WINMEM32.DLL which comes with the Software Development Kit (SDK) provides a set of functions that allow an application to make use of the 32-bit capabilities of the 80386 processor. It contains a function Global32Alloc that allocates a USE32 data segment (up to 16Mbytes) for which a code alias can be made with the function Global32CodeAlias, thus enabling segment aliasing. This makes it possible to generate and run 32-bit ML code within Windows.

As mentioned above (parts of) the runtime system should lie in the same address space as the ML code. We shall somehow move the runtime system into the ML-heap allocated with the Global32Alloc function. This can be done by collecting the relevant variables and functions of the runtime system in an assembler module, and then compiling and linking this module into a segment that can be copied into the ML-heap when initializing the system (the Microsoft C compiler cannot generate 32-bit code so we have to use the assembler). It is possible to access an USE32 segment from C code through a 64K "window" obtained by the Global16PointerAlloc function in the WINMEM32 library. This function allocates an USE16 alias to a portion (up to 64K) of the USE32 segment, but it is inconvenient and slow to use this technique when operating on larger portion of the ML heap. With the assembler one has completely control over and access to all the different kinds of segments.

There are three types of functions in the runtime system:

- functions that are called directly from the compiled ML-code must be placed in the ML-heap and therefore must be implemented in assembler.
- functions that interact closely with the ML-heap; these could be implemented in C by using USE16 aliases to access the ML heap, but have been implemented in assembler for greater efficiency.
- functions that hardly interact with the ML-heap; these can without any loss of efficiency be implemented in C.

The heart of the garbage collector is implemented in assembler together with some few utility functions (as are the functions that are called directly from ML). The rest of the runtime system is implemented in C. Access to the ML-heap is obtained through 64K "windows" allocated with Global16PointerAlloc function.

3 The runtime system

In this section the organization of the new runtime system is described in terms of the existing SML.NJ runtime systems.

The SML.NJ runtime system is written in C and assembler. It consists of:

- A garbage collector
- Small library functions written in assembler
- Small library functions written in C
- Facilities to handle Export and Import
- Facilities to handle signals (interrupts)

The runtime systems for the different UNIX versions are very alike. The differences are mostly expressed using #IFDEF OPSYS ... #ENDIF declarations in
3 THE RUNTIME SYSTEM

the C-code. When writing a runtime system for Windows we cannot simply reuse the original C-code. As mentioned in the previous section, there are problems with mixing USE16 C-code and USE32 ML-code. However, I have tried to maintain the original structure of the runtime system, and only made modifications where needed because of the mixed memory model and the restrictions made by Windows. (Before reading the C code of the new runtime system, the reader is advised first to look at the original runtime system written for UNIX, because the many non-standard C details in the Windows version reduce the clarity.)

3.1 Segments

One of the first things the system does, when started, is to setup the USE32 segments that constitute the ML-heap. This is done by the code in the file segments.c. A USE32 segment (called Use32Data) is allocated and a code alias (called Use32Code) is established. These two constitute the ML-heap. The part of the runtime system that must lie on the ML-heap are placed in an assembler module (in the file prim.asm) which is compiled and linked into a segment (called .RUNCODE). The contents of this segment is copied into the ML-heap by the code that set up the USE32 segments.

3.2 The interface between ML and the runtime system

After the system is initialized, control is passed to the compiled ML code. A C structure which contains the addresses of relevant variables and functions in the runtime system is passed on to the ML code. To call a C library function the ML code looks up its address faddr in a table and calls the callC function with faddr as an argument.

Our interface must handle the context switch from USE16 to USE32 associated with the transfer of control from the runtime system (USE16) to the compiled ML code (USE32). When the ML code is entered, registers that the C compiler uses are saved on the stack. A C structure (MLState) holds the ML state. Before jumping to ML, registers are loaded from this structure. When returning to the runtime system things happens in reverse order.

When executing code in the runtime system the values in the segment registers are taken care of by the C compiler/linker and the Windows loader. Before jumping to the compiled ML-code we must load the segment registers with the correct values. DS is used as implicit segment register in most machine instructions. ES is used in string operations, and the SS segment register is used when ESP and EBP serve as index registers in memory references. DS, ES, and SS are loaded with the Use32Data value returned by GlobalUse32Alloc, before jumping to the ML-code. Care must be taken when switching stack in this way. When the USE16 stack is used, the upper 16 bits of the stack pointer (ESP) are not used, and can have random values. When switching to the USE32 stack we must ensure that the ESP register contains a legal stack address. After a move to the SS register the 80386 interrupt is disabled in one instruction, so we can handle this by the following code sequence:

```
...  
mov ax, Use32Data
mov ss, ax ; no interrupts between
```
mov esp, Use32StackPointer ; these two instructions
...

The context switch is handled by the function restoreregs in the file interface.asm and by the functions enterUse32 and saveregs in the file prim.asm, which also contains the variables and functions which must be placed in the ML-heap.

### 3.3 C library functions

The UNIX runtime system contains a number of C library functions that handle the interaction with the operating system. Some of these are UNIX specific and have no equivalent in Windows. In the Windows runtime system, these will generate a system exception if called:

```c
void foo(ML_val_t arg)
{ raise_syserror("foo is not implemented under Windows!"); }
```

Certain functions that are not crucial to the compiler and that are difficult to implement under Windows are not implemented yet, and will also generate a system exception, if called.

The functions crucial to the compiler are all implemented. Some of these functions are UNIX specific and do not have an equivalent meaning in Windows, e.g. the masking and unmasking of signals. When called, those functions will perform some neutral action but will not cause a system exception. This will in some cases mean wrong return values to the ML system but these are not crucial. For example, Windows cannot perform all of the timer functions that are available in UNIX, so timing is not accurate in the Windows system. The C library functions are located in the file callc.c.

Details about the interaction between the runtime system and the ML code and data are found in [5], and in the comments in the files prim.asm, gc.asm, interface.asm, and util.asm.

### 3.4 Assembly library functions

There are a number of assembler library functions that are called directly from compiled ML code. These are:

- `array(n,x)` allocates an array of length `n`, and initializes its elements to `x`.
- `callc(f,a)` calls the C function whose address is `f` with the argument `a`.
- `create_b(n)` allocates an uninitialized byte-array of length `n`.
- `create_s(n)` allocates a string of length `n`.
- `floor(x)` returns the "floor" of the real number `x`.
- `logb(x)` returns the exponent of the real number `x`.
- `scalb(x)` inserts a new exponent into the real number `x`.

Floating point operations are executed by using the 80387 math co-processor. We have added utility functions to handle the mixed memory model. See comments in the file util.asm.
3 THE RUNTIME SYSTEM

3.5 Garbage collection

The original garbage collector operates on the entire ML heap. Because it is difficult and inefficient to access USE32 data from the C code, we have implemented the heart of the garbage collector in assembler. It is actually just a "hand compilation" of the original garbage collector written in C, where 32-bit addresses and registers are used. The heart of the garbage collector is located in the file gc.asm, whereas other g.c. related functions are found in the file callgc.c.

3.6 Signal handling

In the UNIX version signal handlers have been implemented for a number of hardware signals. This is not possible in the Windows version. The whole signal machinery has been neutralized in the Windows version. The lack of signal handlers has some annoying consequences. For example it is not possible to stop an infinite loop with Ctrl-C.

As shown in [1] signals can, in a smooth way, be used to initiate garbage collection. All the existing implementations for UNIX use signals this way, either by allocating ahead until a pagefault occur, or by an explicit test for the available memory followed by an "interrupt on overflow". This can't be done in the Windows version. Instead we must make an explicit test and jump to a routine that can initiate the garbage collection when necessary.

```
...  
    cmp  datalimit, allocptr  |  
    jno  no_overflow  |  8 bytes  
    jmp  initiate_gc  |  
  no_overflow: ...  
```

instead of:

```
...  
    cmp  datalimit, allocptr  |  
    into  |  4 bytes  
    ...  
```

This is unpleasant because it takes up space (we shall perform this check at the beginning of every function).

3.7 Export and Import

In the UNIX version it is possible to export the state to an executable file (a.out format). This is used to make stand alone programs, and in particular it is used when bootstrapping the system to make "ready to run" versions of the interactive system and the batch compiler. Because of the problems that Windows has in handling USE32 segment this is not possible in Windows.

The ability to export the state to an executable file, has influence on the execution speed and the memory requirements. If exported to an executable file, the compiler code is located together with the runtime system below the base address of the ML heap, and will therefore not be collected when a major collection is performed [1]. This would in our system mean higher execution speed because most of
4 THE CODE GENERATOR

functor CPSgen( Machine : CMACHINE ) =

... fun gen cexp =
    (* generate code for the CPS-expression cexp *)
    Machine.subl3t(immed 1, regbind v, arithtemp)
... end

Figure 1: CPSgen converts CPS-expressions to machine code. Functions in the structure which CPSgen is applied to, shall generate the corresponding machine code, when called for by CPSgen.

the time is spent doing major collections. Because the heap always is at least three times the size of the living data [1, 2], the presence of the whole compiler on the heap contributes considerably to the memory requirements.

In principle, we could simulate the export by writing the contents of the ML-heap together with the contents of the runtime system's data segment to a file, and then use a special version of the runtime system to load and execute this file. This has not been done yet; instead the runtime system can load *.mo files as described in the howto-boot file in the SML.NJ documentation directory.

4 The code generator

The SML.NJ is nicely divided into a machine-independent front end and a machine-dependent back end. A single signature (called CMACHINE) defines the interface between the two. The back end consists of a few structures that — together with a few structures in the front end — are described below. At the end of this section some special aspects of the 80386 that has influence on the code generator are examined more closely. The intention with this section is to give a survey of the code generator and the description is therefore kept at a very high level.

4.1 From CPS-expressions to machine code

The front end transforms SML source code into CPS-expressions [3, 5, 7, 8]. The transformation of CPS-expressions into machine code is handled by the machine-independent functor CPSgen. CPSgen is parameterized on elementary code generating functions specified by the signature CMACHINE. That is, CMACHINE defines names for datatypes, variables, and functions that the front end will use when transforming CPS-expressions into machine code (see Figure 1). The CPSgen functor is located in the file generic.sml, and CMACHINE is found in cmachine.sig.

4.2 Backpatch and jumpsize-optimization

The back end has to handle relative addresses. That is, we have to backpatch relative jumps and other instructions that use relative addressing. To help in this, a machine-independent functor, Backpatch, is included in the compiler. Backpatch is
parameterized on a machine-dependent structure Jump which contains the information needed to backpatch on a particular machine. Backpatch needs to know the size in bytes of the instructions that use relative addressing and how to emit code for these. The signatures BACKPATCH and JUMP are given in Figure 2.

The machine code generator can make use of the primitives named in BACKPATCH (emitstring, newlabel, etc) when implementing the functions in CAMCHINE. For example the emitstring() is used every time a string (code or data) is put into the code (see Figure 3). The full Backpatch functor, which is part of the general SML NJ system, can be found in the file backpatch.sml. The Jump structure for the 80386 is found in the file 386jumps.sml.

4.3 The 80386 code generator

As mentioned in section 4.1 we have to write a structure that matches the CAMCHINE signature. A functor CMach386 is used for this. The front end includes a batch compiler that can generate assembler code, so we need to generate both assembly code and machine code. The 'argument' given to CMach386 determines which one is generated.

The signature CODER386 defines a subset of the 80386 instruction set and addressing modes, that is used in the code generation. Two functors which match this signature

MCode386: generates machine code
ACode386: generates assembly code

are outlined in Figure 3.

We see how the functor MCode386 is parameterized on the machine-dependent Jump structure mentioned above, and how this structure is passed on to Backpatch. This is how we get access to the primitives named in the BACKPATCH signature. When generating assembly code we do not need to backpatch and therefore the Jump structure is unnecessary. Instead symbolic labels are used. Notice that the primitives in Backpatch are used in the MCode386 module only.

These two functors are used to make a 80386 code generator to be passed on to CPSgen. Applying the functor CMach386 to one of them results in a structure whose functions when called, generates code (machine or assembly) for the 80386.

The signature CODER386 is found in the file 386coder.sig. The Mcode386 and the ACode386 functors are found in the files 386mcode.sml and 386acode.sml.

4.4 Putting it together

SML NJ includes an interactive system and a batchcompiler. The front end functor IntShare defines the interactive system, and Batch defines the batchcompiler. Figure 4 shows how to build the interactive system and the batchcompiler for the 80386. Batch takes 2 arguments; M that generates machine code and A that generates assembly code. Intshare takes three argument where one of them (Machm) is the structure that generates the machine code. Comp386 is the batchcompiler module and Int386 is the interactive module.
signature BACKPATCH =
    (* Machine-independent *)
    sig
        eqtype Label
        type JumpKind
        (* Note 1 *)
        val newlabel : unit -> Label (* Create a new label *)
        val define : Label -> unit (* Associate a label with a point in the code *)
        val emitstring: string -> unit (* Insert a string into the code *)
        val align : unit -> unit (* Ensure that the next code is on a 4-byte boundary *)
        val jump : JumpKind*Label -> unit (* Insert a JumpKind instruction into the code (note 2) *)
        val mark : unit -> unit (* Insert a gc-tag in the code (note 3) *)
        val finish : unit -> string (* Initiate the backpatching. *)
    end

signature JUMPS =
    (* Machine-dependent *)
    sig
        type JumpKind
        val sizejump : JumpKind*int*int*int -> int (* return the size of the JumpKind instruction *)
        val emitjump : JumpKind*int*int*int -> string (* emit code for the JumpKind instruction (note 1) *)
        val emitlong : int -> string (* insert a 32-bit litteral in the code *)
    end

functor Backpatch(Jump : JUMPS) : BACKPATCH =
    (* Machine-independent *)
    struct
        open Jump
        ...
        datatype Desc = ...
            | JUMP of Jumpkind * Label * int ref * desc | ...
        ...
        fun jump(k,lab) = refs := JUMP(k,lab,ref 0, !refs)
        ...
    end (* functor Backpatch *)

Note 1: The JumpKind datatype is used to encode the different instructions that uses relative addresses. The fragment of backpatch shown here indicates how the "jumps" are inserted into a tree to be backpatched later.

Note 2: The code generator calls this function when instructed to generate code for an instruction that refers to a relative address (i.e. to labels). jump makes a note of this instruction, and later when the backpatching in initiated, Backpatch can determine the size of the instruction (using sizejump) and how to emit code for it (using emitjump).

Note 3: Garbage collection (GC) is explained in [1].

Figure 2: The BACKPATCH and JUMP signatures
(* The basic machine code generation *)
functor MCode386(Jumps : JUMPS386) : CODER386 =
struct
  structure Emitter : BACKPATCH = Backpatch(Jumps)
  open Jumps Emitter
  datatype EA = ...
  fun movl(x : EA, y : EA) = emitstring (... (* build up the string that constitute a move instruction *))
  ...
  (* notice how the emitstring from Backpatch is used *)
end (* functor MCode386 * )

(* The basic assembly code generation *)
functor ACode386() : CODER386 =
struct
  datatype EA = ...
  fun movl(x El, y El) = (emit "mov "; emit2args(x,y))
  ...
end (* functor ACode386 * )

(* The abstract machine *)
functor CMach386( Coder : CODER386) : CMACHINE =
struct
  datatype EA = ...
  fun move(x : EA, y : EA) = Coder.movl(x,y)
  ...
end (* functor CMach386 * )

Note: When CPSgen calls the abstract instruction move(); one of the movl() functions will be called to generate the desired code.

Figure 3: The basic code generators for the 80386 (machine and assembly code).
structure MC386 : CODEGENERATOR =
struct
    structure MachineCoder = MCode386(Jumps386)
    structure CMachine = CMach386(MachineCoder)
    structure MachineGen = CPScomp(CMachine)

    fun generate lexp = (MachineGen.compile lexp; MachineCoder.finish())
end (* structure MC386 *)

structure AC386 : ASSEMBLER =
struct
    structure AssemCoder = ACode386()
    structure CMachine = CMach386(AssemCoder)
    structure AssemGen = CPScomp(CMachine)

    fun generate(lexp,stream) = (AssemGen.compile lexp)
end (* structure AC386 *)

structure Int386 = IntShare(structure Machm = MC386
    val fileExtension = ".386"
    structure D = BogusDbg)

structure Comp386 = Batch(structure M=MC386 and A=AC386)

Figure 4: The 386 glue
4 THE CODE GENERATOR

4.5 Special conditions in 80386

The compiler was originally designed for the Vax and M680x0 family. Later on, code generators have been made for other architectures like MIPS and SPARC. The 80386 lacks some of their facilities and this has caused some problems.

One problem is the number of general registers. The CMACHINE signature specifies a set of variables. These are supposed to be in registers. But the 80386 has only seven general registers, so we have to simulate some extra registers in the memory. Where should these be placed? Under Windows it is possible to use absolute addresses because we manually allocate a segment to hold the ML code and data, and therefore have control over what goes where. Thus we can allocate say the 4n lowest addresses in the ML-heap to hold the n "memory registers". But in general (i.e. in other operating systems) we cannot use absolute addresses.

The runtime system (written in C) defines a set of variables that compiled ML code uses. These (or their addresses) are placed in a C-structure whose address is passed on to the ML code when initializing the system. In the same way we could allocate space in the runtime system (e.g. int MemRegs[n]) and pass MemRegs to the ML code. But this would require some changes in the front end, and the goal was to implement the code generator without any changes at all in the front end.

As a result of Continuation Passing Style, ML does not use the stack. We therefore have a static stack when running ML code, so the "memory registers" can be put on the stack in the same way as one would allocate space for local variables on the stack in a C-compiler. This is what has been done here. But care must be taken when referring to these variables. Some registers in the 80386 are dedicated to special purposes in some instructions. Consequently, the stack is used by a few functions in the code generator to save the value of these registers, when necessary. But this is not a problem because we always know exactly how much has been pushed onto the stack.

Using the stack to hold the memory registers has another advantage over absolute addresses in that we save 2 bytes in the code size on every reference to memory variables. The code size is a serious problem when running under Windows because we are restricted to 16 Mbyte.

Because the 80386 cannot perform memory to memory operations, a lot of moving to temporary registers takes place, which again contributes to the code size. This is a serious problem with the 80386 when running under Windows.

Another problem with the Intel 80386 microprocessor is that it does not directly support PC-relative addressing. SML_NJ provides facilities to handle this problem. The front end keeps track of the addressing requirements of each function and calls the beginStdFn in CMACHINE if it uses PC-relative addressing. One of the parameters to beginStdFn is the closure of the function so beginStdFn can load the base address and later use it for relative addressing (see the file cmachine.sig for details). But in version 0.66 of the compiler, the needPC function (located in CPSgen) that determine whether a function uses PC-relative addressing is constantly true:

\[ \text{fun needPC cexp = true} \]

Thus, if the above technique is used, every function gets the extra overhead with calculating a function's base address. Hoping that many functions don't use PC-relative addressing I have chosen another technique. We need to generate code for instructions like load effective address:
5 Example

In this section we show how a simple function is compiled with the 80386 code generator. The assembly code shown below is generated with the assemble command in the batch compiler (see the file BATCHINSTALL).

\[ \text{fun } f \ x = \begin{cases} 1 & \text{if } x = 0 \\ x \cdot f(x - 1) & \text{else} \end{cases} \]

is compiled into:

\[
\begin{align*}
& \text{L3:} \\
& \text{cmp} \quad \text{dword ptr} \ [\text{esp}+0], \text{edi} \quad ; \text{check the available memory} \\
& \text{jns} \ \text{02} \\
& \text{call} \quad \text{dword ptr} \ [\text{esp}+12] \quad ; \text{and call garbage collection if necessary} \\
& \text{02:} \\
& \text{cmp} \quad \text{dword ptr} \ [\text{esp}+40], 1 \quad ; \text{if } x = 0 \\
& \text{jne} \ \text{L14} \\
& \text{mov} \quad \text{edx}, \text{dword ptr} \ [\text{ebp}+0] \quad ; \text{then continue with } 1 \\
& \text{mov} \quad \text{dword ptr} \ [\text{esp}+44], \text{ebp} \\
& \text{mov} \quad \text{dword ptr} \ [\text{esp}+40], 3 \\
& \text{jmp} \quad \text{edx} \\
& \text{L14:} \quad ; \text{else } x \cdot f(x-1) \\
& \text{mov} \quad \text{eax}, 49 \\
& \text{stos} \quad \text{eax} \\
& \text{lea} \quad \text{eax}, \text{L4} \quad ; \text{setup a closure with the argument } x \text{ and} \\
& \text{stos} \quad \text{eax} \\
& \text{mov} \quad \text{eax}, \text{dword ptr} \ [\text{esp}+40] \quad ; \text{the continuation for the multiplication L4}
\end{align*}
\]
6 Conclusion

We have succeeded in porting the SML.NJ version 0.66 to a PC running Windows. A general 80386 code generator is made, which can be used in other architectures based on the 80386 processor. The new runtime system lacks some of the facilities found in the UNIX version, but we will not continue working on the runtime system until new 32-bit versions of Windows and C compilers becomes available. The purpose of this project was to see if it was possible to implement the whole SML.NJ system on a PC, and that we have proved.

Notice how the stack is used to simulate registers, and how the EAX and ECX are used as temporary registers. In SML.NJ an integer i is represented as i * 2 + 1, so the integer one (1) is represented as the integer three (3).
REFERENCES

No changes have been made in the front end\(^2\), so the whole system is implemented. Other compilers for a subset of ML are available on the PC, but this is the first time (to our knowledge) that a complete SML compiler runs on the PC.

References


\(^2\)We had to change some structure names because filenames in DOS can only be upto 8 characters in length.
Completely Bounded Quantification is Decidable

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Abstract

This paper proves the decidability of subtyping for $F_\leq$ when the bounds on polymorphic types do not contain Top (i.e., in all types of the form $\forall \alpha <.: \tau_1, \tau_2, \tau_1$ does not contain Top). This general restriction is subsequently relaxed to allow unbounded quantification.

1 Introduction

$F_\leq$ [CW85,CG] is a typed $\lambda$-calculus with subtyping and bounded second-order polymorphism. The importance of $F_\leq$ in programming language design is that it provides a simple context for studying the typing problems that arise when subtyping and bounded quantification are added to polymorphic languages such as ML.

Curien and Ghelli [CG] recently developed a subtyping algorithm for $F_\leq$ and proved its partial correctness. Subsequently, Ghelli [Ghe90] presented a termination proof for this algorithm. A mistake was discovered in this termination proof, following which Pierce [Pie92] presented a proof showing that the subtyping problem for general $F_\leq$ types is undecidable.

This paper shows how one can make the subtyping problem decidable by imposing some restrictions on $F_\leq$ types. We first prove the termination of Curien and Ghelli's algorithm when the bounds on all polymorphic types involved do not contain Top. i.e., In all types of the form $\forall \alpha <.: \tau_1, \tau_2, \tau_1$ does not contain Top. Such a bound completely determines the structure of $\alpha$, hence we refer to this as "completely bounded". We later show that this restriction can be relaxed to allow unbounded quantification. Adding records and unions to our system causes it to become undecidable. We are currently working on defining subset restrictions for records and unions similar to those presented in this paper to make subtype checking decidable in the presence of these types.

We are in the process of designing a type system based on $F_\leq$ for a prototyping language called Rapide [BL90,MMM91], and are implementing a subtyping algorithm for this type system. Given the undecidability of subtyping for general $F_\leq$ types, we need to restrict our type system so as to

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make subtyping decidable. Results such as those presented in this paper will aid in determining the necessary restrictions.

In Section 2, we present Curien and Ghelli's algorithm. We prove the termination of this algorithm for completely bounded quantification in Section 3. Section 4 shows how we can relax our restrictions to allow unbounded quantification. Section 5 presents examples of records and unions that cannot be handled by simple extensions of Curien and Ghelli's algorithm. Section 6 concludes the paper by describing plans for future work.

2 Curien and Ghelli's algorithm

A $F _{<_}$ type $\tau$ is either a simple type (such as $Int$), a variable, a function type $(\tau_1 \rightarrow \tau_2)$, or a polymorphic type $(\forall \alpha < : \tau _1 . \tau _2)$. Given a list of assumptions $\Gamma$ and two types $\sigma$ and $\tau$, the subtyping problem is to determine whether or not $\Gamma \vdash \sigma < : \tau$ — i.e., whether or not $\sigma$ is a subtype of $\tau$ given the assumptions in $\Gamma$. The assumptions in $\Gamma$ will all be of the form $\alpha < : \tau$, where $\alpha$ is a type variable and $\tau$ is a type. (This convention — that $\alpha$'s refer to type variables and $\sigma$'s and $\tau$'s refer to types — is used for the rest of the paper. We shall also use $\alpha$ to refer to simple types — the context will make it clear whether a particular $\alpha$ is a variable or a simple type.) $\tau$ in this case is called the bound of $\alpha$ and is referred to as $\Gamma (\alpha)$. Free variables in $\tau$ may only be bounded in other assumptions in $\Gamma$ to the left (earlier in the list) of the assumption containing $\tau$.

Curien and Ghelli's algorithm is presented as a list of axiom schemas and inference rules. These schemas and rules contain templates of subtyping problems. The algorithm proceeds by applying the inference rules backwards to the subtyping problem. If the subtyping problem matches the template below the line of an inference rule, it reduces to subtyping problems that can be derived from the templates above the line of the inference rule. If the subtyping problem matches the template of an axiom schema, the algorithm reports success. In all other cases, the algorithm reports failure.

The axiom schemas and inference rules are listed below:

(NTop) $\Gamma \vdash \sigma < : Top$

(NRefl) $\Gamma \vdash \alpha < : \alpha$

(NVar) $\Gamma \vdash \Gamma (\alpha) < : \tau$

$$\frac{}{\Gamma \vdash \alpha < : \tau}$$

(NArrow) $\Gamma \vdash \tau_1 < : \sigma_1 \quad \Gamma \vdash \sigma_2 < : \tau_2$

$$\frac{}{\Gamma \vdash \sigma_1 \rightarrow \sigma_2 < : \tau_1 \rightarrow \tau_2}$$

(NAll) $\Gamma \vdash \tau_1 < : \sigma_1 \quad \Gamma, \alpha : \tau_1 \vdash \sigma_2 < : \tau_2$

$$\frac{}{\Gamma \vdash \forall \alpha < : \sigma_1, \sigma_2 < : \forall \alpha < : \tau_1 . \tau_2}$$
3 Proof of termination

We prove termination by defining a complexity metric which is finite and positive for each subtyping problem, and show that the application of inference rules causes the complexity of the new subtyping problems generated to decrease. Since the complexity cannot decrease indefinitely, the algorithm will have to terminate.

In 3.1, we define the complexity metric. In 3.2, we show how the complexity metric is affected by applying the various inference rules. We conclude with a condition (Theorem 1) that, if satisfied, will guarantee the termination of the subtyping algorithm. Finally, in 3.3, we show how our restrictions on bounds of polymorphic types satisfies the condition of Theorem 1.

3.1 The complexity metric

We define the size of a type $\tau$ with respect to a list of assumptions $\Gamma$, and refer to this as $size(\tau)_\Gamma$. The complexity of a subtyping problem $\Gamma \vdash \sigma < : \tau$ is defined as:

$$complexity(\Gamma \vdash \sigma < : \tau) \overset{\text{def}}{=} size(\sigma)_\Gamma + size(\tau)_\Gamma$$

$size(\tau)_\Gamma$ is determined by recursively replacing variables in $\tau$ with their respective bounds and then computing the textual size of the resulting expression. $size(\tau)_\Gamma$ is formally defined as:

\[
\begin{align*}
size(\text{Top})_\Gamma &= 1 \\
size(\alpha)_\Gamma &= \begin{cases} 
size(\Gamma(\alpha))_\Gamma & \text{if } \Gamma(\alpha) \text{ is defined} \\
1 & \text{otherwise}
\end{cases} \\
size(\tau_1 \rightarrow \tau_2)_\Gamma &= size(\tau_1)_\Gamma + size(\tau_2)_\Gamma \\
size(\forall \alpha < : \tau_1 . \tau_2)_\Gamma &= size(\tau_1)_\Gamma + size(\tau_2)_{\alpha < : \tau_2}
\end{align*}
\]

Examples:

1. $size(\text{Top} \rightarrow \alpha)_\alpha < : \text{Int} \rightarrow \text{Int}$
   
   $= size(\text{Top})_\alpha < : \text{Int} \rightarrow \text{Int} + size(\alpha)_\alpha < : \text{Int} \rightarrow \text{Int}$
   
   $= 1 + size(\text{Int} \rightarrow \text{Int})_\alpha < : \text{Int} \rightarrow \text{Int}$
   
   $= 1 + size(\text{Int})_\alpha < : \text{Int} \rightarrow \text{Int} + size(\text{Int})_\alpha < : \text{Int} \rightarrow \text{Int}$
   
   $= 1 + 1 + 1 = 3$.

2. $size(\forall \alpha_1 < : (\alpha_2 \rightarrow \text{Int}) . \alpha_1)_\alpha_2 < : \text{Int} \rightarrow \text{Int}$

   $= size(\alpha_2 \rightarrow \text{Int})_{\alpha_2 < : \text{Int} \rightarrow \text{Int}} + size(\alpha_1)_{\alpha_2 < : \text{Int} \rightarrow \text{Int}, \alpha_1 < : \alpha_2 \rightarrow \text{Int}}$

   $= size(\alpha_2)_{\alpha_2 < : \text{Int} \rightarrow \text{Int}} + size(\text{Int})_{...} + size(\alpha_2 \rightarrow \text{Int})_{\alpha_2 < : \text{Int} \rightarrow \text{Int}, \alpha_1 < : \alpha_2 \rightarrow \text{Int}}$

   $= size(\text{Int} \rightarrow \text{Int})_{...} + 1 + size(\alpha_2)_{\alpha_2 < : \text{Int} \rightarrow \text{Int}, \alpha_1 < : \alpha_2 \rightarrow \text{Int}} + size(\text{Int})_{...}$

   $= 2 + 1 + size(\text{Int} \rightarrow \text{Int})_{...} + 1$

   $= 2 + 1 + 2 + 1 = 6$. 

Lemma 1 \( \text{size}(\tau) \) is always finite and positive.

Proof. It is obvious from the definition of \( \text{size} \) that it has to be positive. We prove that it is finite by showing that evaluation of \( \text{size} \) terminates for all \( \tau \) and \( \Gamma \). A complexity metric similar to that used in Ghelli's flawed termination proof actually works in this case.

The complexity metric to prove the termination of the evaluation of \( \text{size}(\tau) \) is obtained by first ordering all the variables that occur in \( \tau \) and \( \Gamma \) such that the following property is satisfied: If \( \alpha_i \) is defined in the bound of \( \alpha_j \), then \( \alpha_i \) occurs to the left of \( \alpha_j \) in the ordering. It is possible to obtain such an ordering given the structure of \( F \) (there may be multiple orderings that satisfy this condition in which case, one of them is chosen arbitrarily). The depth of each variable is then defined as the number of variables that occur to the left of it in this ordering.

The complexity of any subproblem \( \text{size}(\tau') \) that arises during the evaluation of \( \text{size}(\tau) \) is the tuple \( (D, S) \), where \( D \) is the maximum depth over all the variables that occur in \( \tau' \) and \( S \) is the textual length of \( \tau' \). There may be variables in \( \tau' \) and \( \Gamma' \) that do not occur in either of \( \tau \) or \( \Gamma \). These variables are created when an existing variable is duplicated in the reduction process, thus requiring one of the uses to be renamed. The depth of the renamed variable is defined to be the same as its unrenamed counterpart. The key to this proof is that defining the depth of renamed variables in this manner maintains the condition based on which the initial ordering was created.

It is easy to see that the complexity decreases (the ordering between \( (D, S) \) tuples is lexicographic) during the evaluation of \( \text{size}(\tau) \). For all reductions other than \( \text{size}(\alpha) = \text{size}(\Gamma(\alpha)) \), the \( D \) component of the complexity metric either remains the same or decreases while the \( S \) component decreases; whereas in the abovementioned reduction, the \( D \) component decreases.

Since this complexity metric cannot decrease indefinitely, the evaluation of \( \text{size} \) for any \( \tau \) and \( \Gamma \) has to terminate.

3.2 The effect of inference rule application on complexity

\( \text{NVar} \)

\[ \text{complexity}(\Gamma \vdash \alpha <: \tau) = \text{size}(\alpha) + \text{size}(\tau) = \text{size}(\Gamma(\alpha)) + \text{size}(\tau) = \text{complexity}(\Gamma \vdash \Gamma(\alpha) <: \tau) \]

i.e., The complexity metric remains the same after application of the inference rule \( \text{NVar} \). However, \( \text{NVar} \) may be applied continuously at most as many times as there are variables in \( \Gamma \) before one of the other rules has to be applied. The complexity metric reduces when any of the other rules are applied, so there is no problem.

\( \text{NArrow} \)

It is quite obvious that \( \text{complexity}(\Gamma \vdash \tau_1 <: \sigma_1) \) and \( \text{complexity}(\Gamma \vdash \sigma_2 <: \tau_2) \) are both less than \( \text{complexity}(\Gamma \vdash \sigma_1 \rightarrow \sigma_2 <: \tau_1 \rightarrow \tau_2) \).
NALL

It is quite obvious that $\text{complexity}(\Gamma \vdash \tau_1 <: \sigma_1)$ is less than $\text{complexity}(\Gamma \vdash \forall \alpha < \sigma_1. \sigma_2 <: \forall \alpha < \tau_1. \tau_2)$. We shall now simplify $\text{complexity}(\Gamma \vdash \forall \alpha < \sigma_1. \sigma_2 <: \forall \alpha < \tau_1. \tau_2) - \text{complexity}(\Gamma, \alpha < \tau_1 \vdash \sigma_2 <: \tau_2)$ to determine the conditions under which the complexity metric decreases when reducing to the second rule above the line in NALL. This expression, which must be positive for the complexity metric to decrease, simplifies as follows:

\[
\text{complexity}(\Gamma \vdash \forall \alpha < \sigma_1. \sigma_2 <: \forall \alpha < \tau_1. \tau_2) - \text{complexity}(\Gamma, \alpha < \tau_1 \vdash \sigma_2 <: \tau_2) = (\text{size}(\forall \alpha < \sigma_1. \sigma_2) \Gamma) - (\text{size}(\forall \alpha < \tau_1. \tau_2) \Gamma) - (\text{size}(\sigma_2) \Gamma, \alpha < \tau_1) + (\text{size}(\tau_2) \Gamma, \alpha < \tau_1)
\]

= \text{size}(\sigma_1) \Gamma + \text{size}(\tau_1) \Gamma + (\text{size}(\sigma_2) \Gamma, \alpha < \sigma_1) - (\text{size}(\tau_2) \Gamma, \alpha < \tau_1)

We concentrate on the part $\text{size}(\sigma_2) \Gamma, \alpha < \sigma_1 - \text{size}(\tau_2) \Gamma, \alpha < \tau_1$ from the last line above. If this expression is non-negative, then the overall expression will be positive, and therefore the complexity metric will decrease on the application of NALL. We need to present the following lemma before we can proceed further.

**Lemma 2** For any types $\sigma$, $\tau$, and for any list of assumptions $\Gamma$, such that the variable $\alpha$ is not bounded in any of them, and also does not occur anywhere in $\Gamma$, $\text{size}(\sigma) \Gamma, \alpha < : \tau = \text{size}(\sigma) \Gamma + n(\text{size}(\tau) \Gamma - 1)$, where $n \geq 0$ and depends only on $\sigma$.

The evaluation of $\text{size}(\sigma) \Gamma$ (for any $\Gamma'$) will reduce to zero or more evaluations of the form $\text{size}(\alpha) \Gamma'$ in addition to other reductions. Reductions to the form $\text{size}(\alpha) \Gamma''$ may be either due to occurrences of $\alpha$ in $\sigma$, or occurrences of $\alpha$ in bounds in $\Gamma'$ which are used to replace the variables they bound. If $\Gamma'$ does not contain any bound that contains $\alpha$, then the number of times $\text{size}(\sigma) \Gamma$ reduces to the form $\text{size}(\alpha) \Gamma''$ depends only on $\sigma$. Suppose this number is $n$.

Therefore, $\text{size}(\sigma) \Gamma$ reduces to $n$ evaluations of the form $\text{size}(\sigma) \Gamma''$, each of which evaluates to 1 since $\alpha$ is not bounded anywhere. So we can write $\text{size}(\sigma) \Gamma$ as $m + n$ where $m$ is the result of the evaluation of the remainder of the reductions.

Similarly, $\text{size}(\sigma) \Gamma, \alpha < : \tau$ reduces to $n$ evaluations of the form $\text{size}(\alpha) \Gamma''$, while the remainder of the reductions evaluate to $m$. Each reduction to $\text{size}(\alpha) \Gamma''$ further reduces to $\text{size}(\tau) \Gamma''$. Since $\Gamma''$ is of the form $\Gamma, \alpha < : \tau, \ldots \tau$ does not depend on the portion of $\Gamma''$ to the right of $\Gamma$. Therefore, $\text{size}(\tau) \Gamma'' = \text{size}(\tau) \Gamma$. So we can write $\text{size}(\sigma) \Gamma, \alpha < : \tau$ as $m + n(\text{size}(\tau) \Gamma)$, which is the same as $\text{size}(\sigma) \Gamma + n(\text{size}(\tau) \Gamma - 1)$. 

We now simplify $\text{size}(\sigma_2) \Gamma, \alpha < : \sigma_1 - \text{size}(\sigma_2) \Gamma, \alpha < : \tau_1$.

\[
\text{size}(\sigma_2) \Gamma, \alpha < : \sigma_1 - \text{size}(\sigma_2) \Gamma, \alpha < : \tau_1 = \text{size}(\sigma_2) \Gamma + n(\text{size}(\sigma_1) \Gamma - 1) - (\text{size}(\sigma_2) \Gamma + n(\text{size}(\tau_1) \Gamma - 1)) = n(\text{size}(\sigma_1) \Gamma - \text{size}(\tau_1) \Gamma)
\]

We are now ready to present the first of our main results.
Theorem 1  During the execution of Curien and Ghelli's algorithm, if size(\(\sigma_1\))\(_{\Gamma} \geq\) size(\(\tau_1\))\(_{\Gamma}\) (where \(\sigma_1\), \(\tau_1\), and \(\Gamma\) are as defined in NALL) is true every time NALL is used to reduce a subtyping problem to the subtyping problem derived from the right template above the line (\(\Gamma, \alpha <: \tau_1 :: \sigma_2 <: \tau_2\)), then the algorithm will terminate.

Proof. Obvious from the results of this section. \(\square\)

3.3 Subset restrictions to guarantee termination

When applying NALL on a subtyping problem, we shall require that we first consider the subtyping problem derived from \(\Gamma \vdash \tau_1 <: \sigma_1\) (the left template above the line in NALL). Only if the algorithm terminates successfully on this problem do we consider the subtyping problem derived from \(\Gamma, \alpha <: \tau_1 :: \sigma_2 <: \tau_2\) (the right template above the line in NALL). Hence, we can assume that \(\Gamma \vdash \tau_1 :: \sigma_1\) when the algorithm is applied on the subtyping problem derived from \(\Gamma, \alpha <: \tau_1 :: \sigma_2 <: \tau_2\). Assuming this, we have the following corollary to Theorem 1.

Corollary 1  For every \(\tau, \sigma\) that are bounds of polymorphic types and for every list of assumptions \(\Gamma\), if \(\Gamma \vdash \tau <: \sigma \Rightarrow\) size(\(\tau\))\(_{\Gamma} \leq\) size(\(\sigma\))\(_{\Gamma}\), then Curien and Ghelli's algorithm will terminate.

Lemma 3  If size(\(\tau_1\))\(_{\Gamma} =\) size(\(\tau_2\))\(_{\Gamma}\), then, for any \(\sigma\), size(\(\sigma\))\(_{\Gamma, \alpha <: \tau_1} =\) size(\(\sigma\))\(_{\Gamma, \alpha <: \tau_2}\).

Proof. This follows trivially from Lemma 2. size(\(\sigma\))\(_{\Gamma, \alpha <: \tau_1} =\) size(\(\sigma\))\(_{\Gamma} + n(size(\(\tau_1\))\(_{\Gamma}\) - 1) =\) size(\(\sigma\))\(_{\Gamma} + n(size(\(\tau_2\))\(_{\Gamma}\) - 1) =\) size(\(\sigma\))\(_{\Gamma, \alpha <: \tau_2}\). \(\square\)

We are now ready to present another key result, Theorem 2, that if the subset restrictions mentioned in Section 1 are satisfied, then the condition of Corollary 1 will be satisfied. A straightforward consequence of this is that if these subset restrictions are met, then Curien and Ghelli's algorithm will terminate, and hence completely bounded quantification is decidable.

Theorem 2  For all types \(\tau, \sigma\) that do not contain Top, and for any list of assumptions \(\Gamma\), if \(\Gamma \vdash \tau <: \sigma\), then size(\(\tau\))\(_{\Gamma} =\) size(\(\sigma\))\(_{\Gamma}\).

Proof. If \(\Gamma \vdash \tau <: \sigma\), then there must be a proof \(\Gamma^1 \vdash \tau^1 :: \sigma^1, \Gamma^2 \vdash \tau^2 :: \sigma^2, \ldots, \Gamma^n \vdash \tau^n :: \sigma^n\) where \(\Gamma^n = \Gamma, \tau^n = \tau, and \sigma^n = \sigma\), and each \(\Gamma^i \vdash \tau^i :: \sigma^i\) is either of the form of NREFL, or obtained from earlier steps in the proof using one of the rules NVAR, NARROW, or NALL. Note that NTop will not be used in such a proof since Top does not occur in \(\sigma\) and \(\tau\).

We prove by induction that for all \(i\) (\(1 \leq i \leq n\)), size(\(\tau^i\))\(_{\Gamma^i} =\) size(\(\sigma^i\))\(_{\Gamma^i}\). The induction hypothesis is that for all \(j\) (\(1 \leq j \leq i\)), size(\(\tau^j\))\(_{\Gamma^j} =\) size(\(\sigma^j\))\(_{\Gamma^j}\). Assuming the induction hypothesis, we prove size(\(\tau^i\))\(_{\Gamma^i} =\) size(\(\sigma^i\))\(_{\Gamma^i}\). There are four cases to consider:

1. \(\Gamma^i \vdash \tau^i :: \sigma^i\) is of the form of NREFL, i.e., \(\tau^i = \sigma^i\). Therefore, size(\(\tau^i\))\(_{\Gamma^i} =\) size(\(\sigma^i\))\(_{\Gamma^i}\).
2. $\Gamma^i \vdash \tau^i <: \sigma^i$ is derived using NVAR from an earlier step of the form $\Gamma^i \vdash \Gamma^i(\tau^i) <: \sigma^i$. In this case, $\tau^i$ is a variable bounded in $\Gamma^i$. Therefore, $\text{size}(\tau^i)_{\Gamma^i} = \text{size}(\Gamma^i(\tau^i))_{\Gamma^i} = \text{size}(\sigma^i)_{\Gamma^i}$.

3. $\Gamma^i \vdash \tau^i <: \sigma^i$ is derived using NARROW from earlier steps $\Gamma^k \vdash \tau^k <: \sigma^k$ and $\Gamma^i \vdash \tau^i <: \sigma^i$. Then $\Gamma^i = \Gamma^k$, $\tau^i = \sigma^k \rightarrow \tau^l$, and $\sigma^i = \tau^k \rightarrow \sigma^i$. Therefore, $\text{size}(\tau^i)_{\Gamma^i} = \text{size}(\sigma^k)_{\Gamma^k} + \text{size}(\tau^l)_{\Gamma^l} = \text{size}(\tau^k)_{\Gamma^k} + \text{size}(\tau^l)_{\Gamma^l} = \text{size}(\tau^k)_{\Gamma^k} + \text{size}(\sigma^i)_{\Gamma^i}$.

4. $\Gamma^i \vdash \tau^i <: \sigma^i$ is derived using NALL from earlier steps $\Gamma^k \vdash \tau^k <: \sigma^k$ and $\Gamma^i \vdash \tau^i <: \sigma^i$. Then $\Gamma^i = \Gamma^k$, $\tau^i = \alpha <: \sigma^k$, $\tau^l = \forall \alpha <: \sigma^k$. Therefore, $\text{size}(\tau^i)_{\Gamma^i} = \text{size}(\sigma^k)_{\Gamma^k} + \text{size}(\tau^l)_{\Gamma^l} = \text{size}(\tau^k)_{\Gamma^k} + \text{size}(\sigma^i)_{\Gamma^i}$.

4 Allowing unbounded quantification

We can relax our restriction on the use of Top to allow unbounded quantification and still retain decidability of subtype checking. Furthermore, any variable bounded by Top (directly or indirectly) may be used as a bound for another variable. We refer to these variables as "Top-bounded".

With this relaxation, there are two kinds of bounds that we can write: (1) Types that do not contain either Top or Top-bounded variables; and (2) Types that are either Top or a Top-bounded variable.

To show that this will not cause problems, we redefine $\text{size}(\text{Top})$ for all Top's that occur as bounds of variables to be a number $L$ that is larger than the size of any bounds of the first kind mentioned above. With this redefinition of size, it is quite easy to see that Corollary 1 continues to hold even when the bounds $\tau$ and $\sigma$ are from this relaxed domain.

There are four cases to consider where $\tau$ and $\sigma$ may each be either of the two kinds mentioned above. We consider each case separately:

1. $\tau$ and $\sigma$ are both types not containing Top or Top-bounded variables: This case has been handled in Theorem 2.

2. $\tau$ and $\sigma$ are both types that are either Top or a Top-bounded variable: Then $\text{size}(\tau) = \text{size}(\sigma) = L$.

3. $\tau$ is a type not containing Top or Top-bounded variables while $\sigma$ is either Top or a Top-bounded variable: Then $\text{size}(\sigma) = L > \text{size}(\tau)$.

4. $\tau$ is either Top or a Top-bounded variable while $\sigma$ is a type not containing Top or Top-bounded variables: In this case, $\tau$ cannot be a subtype of $\sigma$, so we need not consider it any further.
5 Record and union types

We have shown that one can achieve a decidable type system for a fairly unconstricted subset of $F_\leq$. An immediate extension that we started working on was the addition of record and union types. However, it turns out that adding either record or union types to the type system (along with their associated inference rule) makes the subtyping problem undecidable. Finding the right restrictions to allow the addition of these types is the subject of current research.

5.1 Record types

We denote the record type with fields $l_1 \ldots l_n$ having types $\tau_1 \ldots \tau_n$ respectively as $\{l_1: \tau_1, \ldots, l_n: \tau_n\}$.

The inference rule for records is:

\[
\frac{\Gamma \vdash \sigma_1 <: \tau_1 \quad \ldots \quad \Gamma \vdash \sigma_n <: \tau_n}{\Gamma \vdash \{l_1: \sigma_1, \ldots, l_n: \sigma_n, l_{n+1}: \tau_{n+1}, \ldots\} <: \{l_1: \tau_1, \ldots, l_n: \tau_n\}}
\]

(NRec)

Essentially one can either add extra fields to a record type or specialize the types of existing fields to get a subtype.

As we saw earlier, the problem with the original unrestricted system was that one could have subtypes that were structurally much more complicated than the supertype and one could exploit this in creating subtyping subproblems that grew infinitely in their complexity. We prevented this finally by restricting the use of $Top$, which is what allowed a subtype to be more complicated than the supertype in the first place. However, record types provide another way of allowing a subtype to be more complex than the supertype, namely by adding extra fields. So the empty record type provides an entity conceptually similar to $Top$. We now show that one can reproduce the non-terminating example mentioned in Pierce [Pie92], in spite of the restrictions on $Top$ with the use of records.

Let $\neg \tau = \tau \rightarrow b$ where $b$ is some simple type. Note that $\neg \sigma <: \neg \tau$ iff $\tau <: \sigma$.

Let

\[
\theta = \{a: \forall \alpha <: \{\}. \neg\{a: \forall \beta <: \alpha. \neg \beta\}\}
\]

Now consider the subtyping problem

\[
\alpha_0 <: \theta \vdash \alpha_0 <: \{a: \forall \alpha_1 <: \alpha_0. \neg \alpha_1\}
\]

This causes the following sequence of subproblems to be generated infinitely:

\[
\begin{align*}
\alpha_0 &: \theta \vdash \alpha_0 <: \{a: \forall \alpha_1 <: \alpha_0. \neg \alpha_1\} <: \{a: \forall \alpha_1 <: \alpha_0. \neg \alpha_1\} \\
\alpha_0 &: \theta \vdash \{a: \forall \alpha_1 <: \{\}. \neg\{a: \forall \alpha_2 <: \alpha_1. \neg \alpha_2\}\} <: \{a: \forall \alpha_1 <: \alpha_0. \neg \alpha_1\} \\
\alpha_0 &: \theta, \alpha_1 <: \alpha_0 \vdash \neg\{a: \forall \alpha_2 <: \alpha_1. \neg \alpha_2\} <: \neg \alpha_1 \\
\alpha_0 &: \theta, \alpha_1 <: \alpha_0 \vdash \alpha_1 <: \{a: \forall \alpha_2 <: \alpha_1. \neg \alpha_2\} <: \{a: \forall \alpha_2 <: \alpha_1. \neg \alpha_2\} \\
\alpha_0 &: \theta, \alpha_1 <: \alpha_0 \vdash \alpha_0 <: \{a: \forall \alpha_2 <: \{\}. \neg\{a: \forall \alpha_3 <: \alpha_2. \neg \alpha_3\}\} <: \{a: \forall \alpha_2 <: \alpha_1. \neg \alpha_2\} \\
\alpha_0 &: \theta, \alpha_1 <: \alpha_0 \vdash \{a: \forall \alpha_2 <: \{\}. \neg\{a: \forall \alpha_3 <: \alpha_2. \neg \alpha_3\}\} <: \{a: \forall \alpha_2 <: \alpha_1. \neg \alpha_2\} \\
\alpha_0 &: \theta, \alpha_1 <: \alpha_0, \alpha_2 <: \alpha_1 \vdash \neg\{a: \forall \alpha_3 <: \alpha_2. \neg \alpha_3\} <: \neg \alpha_2
\end{align*}
\]
and so on.
This pattern of non-termination is practically identical to the one displayed in [Pie92].

5.2 Union types

Union types pose problems similar to that of record types.
Union types are written as \{t_1: \tau_1 + \ldots + t_n: \tau_n\}, with the usual meaning. The inference rule for union types is fairly straightforward.

\[
\frac{\Gamma \vdash \sigma_1 <: \tau_1 \quad \ldots \quad \Gamma \vdash \sigma_n <: \tau_n}{\Gamma \vdash \{t_1: \sigma_1 + \ldots + t_n: \sigma_n\} <: \{t_1: \tau_1 + \ldots + t_n: \tau_n + \ldots\}}
\]

The subtype is allowed to be a union over a subset of specializations of the types in the supertype.
Since the subtype can be a union of a fewer types than the supertype, one can have types with unions in contravariant positions thereby resulting in subtypes that have more complex structures than the supertype. Again, one can exploit this to generate an example of a subtyping problem that doesn't terminate. Behavior identical to that displayed in the previous example on record types arises for the subtyping problem

\[
\alpha_0 <: \theta \vdash \alpha_0 <: \neg\{(a: \forall \alpha_1 <: \alpha_0. \neg \alpha_1)\}
\]

where

\[
\theta = \neg\{(a: \forall \alpha <: \neg\{\{a: \forall \beta <: \alpha. \neg \beta\}\}\})
\]

6 Future work

We are currently working on various possible subset restrictions on records and unions to make subtype checking decidable in the presence of these types. Following this, we intend to extend our system with recursive types. This might involve techniques similar to those employed by Amadio and Cardelli [AC91] to study the interaction of recursive types with subtyping. A further extension would be the addition of the so-called F-bounds [CCH*89], which essentially allow the bounds of polymorphic functions to be recursive.

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References


An Extension of ML with First-Class Abstract Types

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1 Introduction

Many statically-typed programming languages provide an abstract data type construct, such as the package in Ada, the cluster in CLU, and the module in Modula2. In these languages, an abstract data type consists of two parts, interface and implementation. The implementation consists of one or more representation types and some operations on these types; the interface specifies the names and types of the operations accessible to the user of the abstract data type.

ML [MTH90] provides two distinct constructs for describing abstract data types:

- The (obsolete) abstype mechanism is used to declare an abstract data type with a single implementation. It has been superseded by the module system.
- The ML module system provides signatures, structures, and functors. Signatures act as interfaces of abstract data types and structures as their implementations; functors are essentially parametrized structures. Several structures may share the same signature, and a single structure may satisfy several signatures. However, structures are not first-class values in ML for type-theoretic reasons discussed in [Mac86] [MH88]. This leads to considerable difficulties in a number of practical programming situations.

Mitchell and Plotkin show that abstract types can be given existential type [MP88]. By stating that a value \( v \) has the existential type \( \exists \alpha. \tau \), we mean that for some fixed, unknown type \( \xi \), \( v \) has type \( \tau [\xi/\alpha] \). This paper presents a semantic extension of ML, where the component types of a datatype may be existentially quantified. We show how datatypes over existential types add significant flexibility to the language without even changing ML syntax; in particular, we give examples demonstrating how we express:

- first-class abstract types,
- multiple implementations of a given abstract type,
- heterogeneous aggregates of different implementations of the same abstract type, and
- dynamic dispatching of operations with respect to the implementation type.

We have a deterministic Damas-Milner inference system [DM82] [CDDK86] for our language, which leads to a syntactically sound and complete type reconstruction algorithm. Furthermore, the type system is semantically sound with respect to a standard denotational semantics.

Most previous work on existential types does not consider type reconstruction. Other work appears to be semantically unsound or does not permit polymorphic instantiation of variables of existential type. By contrast, in our system such variables are let-bound and may be instantiated polymorphically.

We have implemented a Standard ML prototype of an interpreter with type reconstruction for our core language, Mini-ML [CDDK86] extended with recursive datatypes over existentially quantified component types. All examples from this paper have been developed and tested using our interpreter.

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2 ML Datatypes with Existential Component Types

In ML, datatype declarations are of the form

```
datatype [arg] T = K_1 of \tau_1 | ... | K_n of \tau_n
```

where the K's are value constructors and the optional prefix argument arg is used for formal type parameters, which may appear free in the component types \( \tau_i \). The value constructor functions are universally quantified over these type parameters, and no other type variables may appear free in the \( \tau_i \)'s.

An example for an ML datatype declaration is

```
datatype 'a Mytype = mycons of 'a * ('a -> int)
```

Without altering the syntax of the datatype declaration, we now give a meaning to type variables that appear free in the component types, but not in the type parameter list. We interpret such type variables as existentially quantified.

For example,

```
datatype Key = key of 'a * ('a -> int)
```

describes a datatype with one value constructor whose arguments are pairs of a value of type 'a and a function from type 'a to int. The question is what we can say about 'a. The answer is, nothing, except that the value is of the same type 'a as the function domain. To illustrate this further, the type of the expression

```
key(3, fn x => 5)
```

is Key, as is the type of the expression

```
key([1, 2, 3], length)
```

where length is the built-in function on lists. Note that no argument types appear in the result type of the expression. On the other hand,

```
key(3, length)
```

is not type-correct, since the type of 3 is different from the domain type of length.

We recognize that Key is an abstract type comprised by a value of some type and an operation on that type yielding an int. It is important to note that values of type Key are first-class; they may be created dynamically and passed around freely as function parameters. The two different values of type Key in the previous examples may be viewed as two different implementations of the same abstract type.

Besides constructing values of datatypes with existential component types, we can decompose them using the let construct. We impose the restriction that no type variable that is existentially quantified in a let expression appears in the result type of this expression or in the type of a global identifier. Analogous restrictions hold for the corresponding open and abstype constructs described in [CW85] [MP88].

For example, assuming x is of type Key, then

```
let val key(v, f) = x in
  f v
end
```

has a well-defined meaning, namely the int result of f applied to v. We know that this application is type-safe because the pattern matching succeeds, since x was constructed using constructor key, and at that time it was enforced that f can safely be applied to v. On the other hand,

```
let val key(v, f) = x in
  v
end
```

is not type-correct, since we do not know the type of v statically and, consequently, cannot assign a type to the whole expression.
Our extension to ML allows us to deal with existential types as described in [CW85] [MP88], with the further improvement that decomposed values of existential type are \texttt{let}-bound and may be instantiated polymorphically. This is illustrated by the following example,

```ml
datatype 'a t
  = k of ('a -> 'b) * ('b -> int)

let val k(f1,f2) = k(fn x => x,fn x => 3) in
  (f2(f1 7),f2(f1 true))
end
```

which results in \((3, 3)\). In most previous work, the value on the right-hand side of the binding would have to be bound and decomposed twice.

3 Some Motivating Examples

Minimum over a heterogeneous list

Extending on the previous example, we first show how we construct heterogeneous lists over different implementations of the same abstract type and define functions that operate uniformly on such heterogeneous lists. A heterogeneous list of values of type \texttt{Key} could be defined as follows:

```ml
val hetlist = [key(3,fn x => x), key([1,2,3,4],length), key(7,fn x => 0),
              key(true,fn x => if x then 1 else 0), key(12,fn x => 3)]
```

The type of \texttt{hetlist} is \texttt{Key list}; it is a homogeneous list of elements each of which could be a different implementation of type \texttt{Key}. We define the function \texttt{min}, which finds the minimum of a list of \texttt{Key}'s with respect to the integer value obtained by applying the second component (the function) to the first component (the value).

```ml
fun min [x] = x
  | min ((key(v1,f1))::xs) = let val key(v2,f2) = min xs in
    if f1 v1 <= f2 v2 then key(v1,f1) else key(v2,f2)
  end
```

Then \texttt{min hetlist} returns \texttt{key (7, fn x => 0)}, the third element of the list.

Stacks parametrized by element type

The previous examples involved datatypes with existential types but without polymorphic type parameters. As an example for a type involving both, we show an abstract stack parametrized by element type.

```ml
datatype 'a Stack = stack of {value : 'b,
                          empty : 'b,
                          push : 'a * 'b -> 'b
                          pop : 'b -> 'a * 'b
                          top : 'b -> 'a,
                          isempty : 'b -> bool}
```

An implementation of an \texttt{int} \texttt{Stack} in terms of the built-in type \texttt{list} can be given as

```ml
stack(value = [1,2,3], empty = [], push = op ::,
      pop = fn xs => (hd xs,tl xs), top = hd, isempty = null)
```

An alternative implementation of \texttt{Stack} could be given, among others, based on arrays. Different implementations could then be combined in a list of stacks. To facilitate dynamic dispatching, constructors of stacks of different implementations can be provided together with stack operations that work uniformly...
across implementations. These “outer” operations work by opening the stack, applying the intended “inner”
operation, and encapsulating the stack again, for example

```haskell
fun makeliststack xs = stack{value = xs, empty = [], push = op ::,
  pop = fn xs => (hd xs, tl xs), top = hd, isempty = null}
fun makearraystack xs = stack{...}
fun push a (stack{value = v, push = pu, empty = e,
  pop = po, top = t, isempty = i}) =
  stack{value = pu(a, v), push = pu, empty = e,
    pop = po, top = t, isempty = i}
map (push 8) [makeliststack [2, 4, 6], makearraystack [3, 5, 7]]
```

4 Type-Theoretical Aspects

A deterministic type inference system for our language is given in the appendix; it leads directly to a syn-
tactically sound and complete type reconstruction algorithm to compute principal types. Our type system is
semantically sound with respect to a standard denotational semantics. Moreover, it is a conservative exten-
sion of ML. That is, for a program in our language whose declarations introduce no existentially quantified
type variables, our type reconstruction algorithm and the ML type reconstruction algorithm compute the
same type. A comprehensive treatment of polymorphic type inference with existential types is found
in [La92].

5 Related Work

Hope+C

The only other work known to us that deals with Damas-Milner-style type inference for existential types
is [Per90]. However, the typing rules given there are not sufficient to guarantee the absence of runtime type
errors, even though the Hope+C compiler seems to impose sufficient restrictions. The following unsafe pro-
gram, here given in ML syntax, is well-typed according to the typing rules, but rejected by the compiler:

```ml
datatype T = K of ''a
fun f x = let val K z = x in z end
f(K 1) = f(K true)
```

XML+

The possibility of making ML structures first-class by implicitly hiding their type components is discussed
in [MMM91] without addressing the issue of type inference. By hiding the type components of a structure,
its type is implicitly coerced from a strong sum type to an existential type. Detailed discussions of sum types
can be found in [Mac86] [MH88].

Haskell with existential types

Existential types combine well with the systematic overloading polymorphism provided by Haskell type
classes [WB89]; this point is further discussed in [LO91]. Briefly, we extend Haskell’s data declaration in a
similar way as the ML datatype declaration above. In Haskell [HPW91], it is possible to specify what type
class a (universally quantified) type variable belongs to. In our extension, we can do the same for existen-
tially quantified type variables. This lets us construct heterogeneous aggregates over a given type class.

Dot notation

MacQueen [Mac86] observes that the use of existential types in connection with an elimination construct
(open, abstype, or our let) is impractical in certain programming situations; often, the scope of the
elimination construct has to be made so large that some of the benefits of abstraction are lost. A formal treat-
ment of the dot notation, an alternative used in actual programming languages, is found in [CL90]. An ex-
tension of ML with an analogous notation is described in [La92].
Dynamics in ML

An extension of ML with objects that carry dynamic type information is described in [LM91]. A dynamic is a pair consisting of a value and the type of the value. Such an object is constructed from a value by applying the constructor `dynamic`. The object can then be dynamically coerced by pattern matching on both the value and the runtime type. Existential types are used to match dynamic values against dynamic patterns with incomplete type information. Dynamics are useful for typing functions such as `eval`. However, they do not provide type abstraction, since they give access to the type of an object at runtime. It seems possible to combine their system with ours, extending their existential patterns to existential types. We are currently investigating this point.

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A Formal Discussion of the Extended Language

In this appendix, we describe the formal language and the type system underlying our extension of ML. The typing rules and auxiliary functions translate to the type reconstruction algorithm given below.

A.1 Syntax

Language syntax

- **Identifiers**: \( x \)
- **Constructors**: \( K \)
- **Expressions**: \( e ::= x | (e_1, e_2) | e | e' | \lambda x.e | \text{let } x = e \text{ in } e' | \text{data } \forall \alpha_1 \ldots \alpha_n. \chi \text{ in } K \mid \text{is } K \mid \text{let } K \ x = e \text{ in } e' \)

In addition to the usual constructs (identifiers, applications, \( \lambda \)-abstractions, and let expressions), we introduce desugared versions of the ML constructs that deal with datatypes. A data declaration defines a new datatype; values of this type are created by applying a constructor \( K \), their tags can be inspected using an is expression, and they can be decomposed by a pattern-matching let expression. The following example shows a desugared definition of ML's list type and the associated length function.

**data** \( \forall \alpha. (\mu \beta. \text{Nil unit } + \text{Cons } \alpha \times \beta) \) in

- let \( \text{length } = \text{fix } \lambda x. \text{length}. \lambda x. \)
  - if (is \( \text{Nil } xs \))
    - 0
  - (let \( \text{Cons } ab = xs \text{ in } + (\text{length}(\text{snd } ab)) \) 1)
  - in
  - \( \text{length } (\text{Cons } (3, \text{Cons } (7, \text{Nil } ()))) \)

Type syntax

- **Type variables**: \( \alpha \)
- **Skolem functions**: \( \kappa \)
Types \[ \tau ::= \text{unit} \mid \text{bool} \mid \alpha \mid \tau_1 \times \tau_2 \mid \tau \rightarrow \tau' \mid \kappa(\tau_1, \ldots, \tau_n) \mid \chi \]

Recursive types \[ \chi ::= \mu \beta. K_1 \eta_1 + \ldots + K_m \eta_m \text{ where } K_i \neq K_j \text{ if } i \neq j \]

Existential types \[ \eta ::= \exists \alpha. \eta \mid \tau \]

Type schemes \[ \sigma ::= \forall \alpha. \sigma \mid \tau \]

Assumptions \[ a ::= \sigma/x \mid \forall \alpha_1 \ldots \alpha_n. \chi/K \]

Our type syntax includes recursive types \( \chi \) and Skolem type constructors \( \kappa \); the latter are used to type identifiers bound by a pattern-matching \texttt{let} whose type is existentially quantified. Explicit existential types arise only as domain types of value constructors. Assumption sets serve two purposes: they map identifiers to type schemes and constructors to the recursive type schemes they belong to. Thus, when we write \( A(K) \), we mean the \( \sigma \) such that \( \sigma = \forall \alpha_1 \ldots \alpha_n \ldots + K \eta + \ldots \). Further, let \( \sum[K\eta] \) stand for sum type contexts such as \( K_1 \eta_1 + \ldots + K_m \eta_m \), where \( K_i = K \) and \( \eta_i = \eta \) for some \( i \).

A.2 Type Inference

Instantiation and generalization of type schemes

\[ \forall \alpha_1 \ldots \alpha_n. \tau \geq \tau' \quad \text{iff there are types } \tau_1, \ldots, \tau_n \text{ such that } \tau' = \tau[\tau_1/\alpha_1, \ldots, \tau_n/\alpha_n] \]

\[ \exists \alpha_1 \ldots \alpha_n. \tau \leq \tau' \quad \text{iff there are types } \tau_1, \ldots, \tau_n \text{ such that } \tau' = \tau[\tau_1/\alpha_1, \ldots, \tau_n/\alpha_n] \]

\[ \text{gen } (A, \tau) = \forall (FV(\tau) \setminus FV(A)) \cdot \tau \]

\[ \text{skolem } (A, \exists \gamma_1 \ldots \gamma_n. \tau) = \tau[\kappa_1(\alpha_1, \ldots, \alpha_k)/\gamma_1] \text{ where } \kappa_1 \ldots \kappa_n \text{ are new Skolem type constructors such that } \{\kappa_1, \ldots, \kappa_n\} \cap FS(A) = \emptyset, \text{ and } \{\alpha_1, \ldots, \alpha_k\} = FV(\exists \gamma_1 \ldots \gamma_n. \tau) \setminus FV(A) \]

The first three auxiliary functions are standard. The function \text{skolem} replaces each existentially quantified variable in a type by a unique type constructor whose actual arguments are those free variables of the type that are not free in the assumption set; this reflects the "maximal" knowledge we have about the type represented by an existentially quantified type variable. In addition to \( FV \), the set of free type variables in a type scheme or assumption set, we use \( FS \), the set of Skolem type constructors that occur in a type scheme or assumption set.

Inference rules for expressions

The first five typing rules are essentially the same as in [CDDK86].

\[
\text{(VAR)} \quad \frac{A(x) \geq \tau}{A \vdash x : \tau}
\]

\[
\text{(PAIR)} \quad \frac{A \vdash e_1 : \tau_1 \quad A \vdash e_2 : \tau_2}{A \vdash (e_1, e_2) : \tau_1 \times \tau_2}
\]
The new rules DATA, CONS, TEST, and PAT are used to type datatype declarations, value constructors, is expressions, and pattern-matching let expressions, respectively.

\[
\sigma = \forall \alpha_1 \ldots \alpha_n. \mu \beta. K_1 \eta_1 + \cdots + K_m \eta_m
\]

\[
FV(\sigma) = \emptyset \quad A[\sigma/K_1, \ldots, \sigma/K_m] \vdash e : \tau
\]

The DATA rule elaborates a declaration of a recursive datatype. It checks that the type scheme is closed and types the expression under the assumption set extended with assumptions about the constructors.

\[
A(K) \geq \mu \beta. \Sigma[K\eta] \quad \eta[\mu \beta. \Sigma[K\eta]/\beta] \leq \tau
\]

The CONS rule observes the fact that existential quantification in argument position means universal quantification over the whole function type; this is expressed by the second premise.

\[
A(K) \geq \mu \beta. \Sigma[K\eta]
\]

The TEST rule ensures that is is K is applied only to arguments whose type is the same as the result type of constructor K.

\[
A \vdash e : \mu \beta. \Sigma[K\eta] \quad FS(\tau') \subseteq FS(A)
\]

The last rule, PAT, governs the typing of pattern-matching let expressions. It requires that the expression e be of the same type as the result type of the constructor K. The body e' is typed under the assumption set extended with an assumption about the bound identifier x. By definition of the function skolem, the new Skolem type constructors do not appear in A; this ensures that they do not appear in the type of any identifier free in e' other than x. It is also guaranteed that the Skolem constructors do not appear in the result type \(\tau'\).

Relation to the ML Type Inference System

Theorem 1 [Conservative extension] Let Mini-ML' be an extension of Mini-ML with recursive datatypes, but not with existential quantification. Then, for any Mini-ML' expression e, \(A \vdash e : \tau\) iff \(A \vdash_{\text{Mini-ML'}} e : \tau\).

Proof: By structural induction on e.

Corollary 2 [Conservative extension] Our type system is a conservative extension of the Mini-ML type sys-
item described in [CDDK86], in the following sense: For any Mini-ML expression \( e \), \( A \vdash e : \tau \) iff \( A \vdash_{\text{Mini-ML}} e : \tau \).

Proof: Follows immediately from the previous theorem.

A.3 Type Reconstruction

The type reconstruction algorithm is a straightforward translation from the deterministic typing rules, using a standard unification algorithm [Rob65] [MM82]. We conjecture that its complexity is the same as that of algorithm \( W \).

Auxiliary functions

In our algorithm, we need to instantiate universally quantified types and generalize existentially quantified types. Both are handled in the same way.

\[
\text{inst}_\forall (\forall \alpha_1 \ldots \alpha_n. \tau) = \tau [\beta_1/\alpha_1, \ldots, \beta_n/\alpha_n] \quad \text{where } \beta_1, \ldots, \beta_n \text{ are fresh type variables}
\]

\[
\text{inst}_\exists (\exists \alpha_1 \ldots \alpha_n. \tau) = \tau [\beta_1/\alpha_1, \ldots, \beta_n/\alpha_n] \quad \text{where } \beta_1, \ldots, \beta_n \text{ are fresh type variables}
\]

The functions \textit{skolem} and \textit{gen} are the same as in the inference rules, with the additional detail that \textit{skolem} always creates fresh Skolem type constructors.

Algorithm

Our type reconstruction function takes an assumption set and an expression, and it returns a substitution and a type expression. There is one case for each typing rule.

\[
\begin{align*}
TC (A, x) &= (\text{Id}, \text{inst}_\forall (A(x))) \\
TC (A, (e_1, e_2)) &= \text{let } (S_1, \tau_1) = TC (A, e_1) \\
&\quad (S_2, \tau_2) = TC (S_1 A, e_2) \\
&\quad \text{in } (S_2 S_1, S_2 \tau_1 \times \tau_2) \\
TC (A, ee') &= \text{let } (S, \tau) = TC (A, e) \\
&\quad (S', \tau') = TC (S A, e') \\
&\quad \beta \text{ be a fresh type variable} \\
&\quad U = \text{mgu} (S' \tau, \tau' \rightarrow \beta) \\
&\quad \text{in } (US'S, U\beta) \\
TC (A, \lambda x. e) &= \text{let } \beta \text{ be a fresh type variable} \\
&\quad (S, \tau) = TC (A [\beta/x], e) \\
&\quad \text{in } (S, S\beta \rightarrow \tau) \\
TC (A, \text{let } x = e \text{ in } e') &= \text{let } (S, \tau) = TC (A, e) \\
&\quad (S', \tau') = TC (S A [\text{gen} (S A, \tau)/x], e') \\
&\quad \text{in } (S'S, \tau')
\end{align*}
\]
\[ TC(A, \text{data } \sigma \text{ in } e) = \text{let } \forall \alpha_1 \ldots \alpha_n, \mu \beta.K_1 \eta_1 + \ldots + K_m \eta_m = \sigma \text{ in} \]
\[ \text{if } FV(\sigma) = \emptyset \text{ then} \]
\[ TC(A[\sigma/K_1, \ldots, \sigma/K_m], e) \]

\[ TC(A, K) = \text{let } \tau = \text{inst}_{\eta}(A(K)) \]
\[ \mu \beta \ldots + K \eta + \ldots = \tau \]
\[ \text{in } (Id, (\text{inst}_2(\eta)) \rightarrow \tau) \]

\[ TC(A, \text{is } K) = \text{let } \tau = \text{inst}_{\eta}(A(K)) \]
\[ \text{in } (Id, \tau \rightarrow \text{bool}) \]

\[ TC(A, \text{let } K x = e \text{ in } e') = \text{let } \bar{\tau} = \text{inst}_{\eta}(A(K)) \]
\[ \mu \beta \ldots + K \eta + \ldots = \bar{\tau} \]
\[ (S, \tau) = TC(A, e) \]
\[ U = (\text{mgu}(\bar{\tau}, \tau)) S \]
\[ \tau_k = \text{skolem}(UA, U\eta) \]
\[ (S', \tau') = TC(UA[\text{gen}(UA, \tau_k)/x], e') \]
\[ \text{in } \]
\[ \text{if } FS(\tau') \subseteq FS(S'UA) \land \]
\[ (FS(\tau_k) \setminus FS(U\eta)) \cap FS(S'UA) \neq \emptyset \]
\[ \text{then } (S'U, \tau') \]

**Theorem 3 [Syntactic Soundness and Completeness]** The type reconstruction algorithm \( TC \) is sound and complete with respect to the type inference relation \( \vdash \).

**Proof:** We extend the proof given in [CDDK86] to deal with the new constructs.

**A.4 Semantics**

We give a standard denotational semantics. The evaluation function \( E \) maps an expression \( e \in \text{Exp} \) to some semantic value \( v \), in the context of an evaluation environment \( p \in \text{Env} \). An evaluation environment is a partial mapping from identifiers to semantic values. Runtime type errors are represented by the special value \( \text{wrong} \). Tagged values are used to capture the semantics of algebraic data types.

We distinguish between the three error situations, runtime type errors (\( \text{wrong} \)), nontermination, and a mismatch when an attempt is made to decompose a tagged value whose tag does not match the tag of the destructor. Both nontermination and mismatch are expressed by \( \bot \).

Our type inference system is sound with respect to the evaluation function; a well-typed program never evaluates to \( \text{wrong} \). The formal proof for semantic soundness is given below.

It should be noted that we do not commit ourselves to a strict or non-strict evaluation function. Therefore, our treatment of existential types applies to languages with both strict and non-strict semantics. For either case, appropriate conditions would have to be added to the definition of the evaluation function.
Semantic domain

Unit value \[ U = \{ \text{unit} \} \]

Boolean values \[ B = \{ \text{false} , \text{true} \} \]

Constructor tags \[ C \]

Semantic domain \[ V = U + B + (V \rightarrow V) + (V \times V) + (C \times V) + \{ \text{wrong} \} \]

In the latter definition of \( V \), + stands for the coalesced sum, so that all types over \( V \) share the same \( \bot \).

Semantics of expressions

The semantic function for expressions, \( E : \text{Exp} \rightarrow \text{Env} \rightarrow V \), is defined as follows:

\[
\begin{align*}
\text{E}[x] \rho &= \rho(x) \\
\text{E}[e_1, e_2] \rho &= \langle \text{E}[e_1] \rho, \text{E}[e_2] \rho \rangle \\
\text{E}[e_1 e_2] \rho &= \text{if } \text{E}[e_1] \rho \in V \rightarrow V \text{ then } \\
&\quad (\text{E}[e_1] \rho)(\text{E}[e_2] \rho) \text{ else wrong} \\
\text{E}[\lambda x.e] \rho &= \lambda v \in V.\text{E}[e](\rho[v/x]) \\
\text{E}[\text{let } x = e \text{ in } e'] \rho &= \text{E}[e'](\rho[\text{E}[e] \rho/x]) \\
\text{E}[\text{data } \sigma \text{ in } e] \rho &= \text{E}[e] \rho \\
\text{E}[K] \rho &= \lambda v \in V.\langle K, v \rangle \\
\text{E}[\text{is } K] \rho &= \lambda v \in V.\text{if } v \in \{ K \} \times V \text{ then true else false} \\
\text{E}[\text{let } K x = e \text{ in } e'] \rho &= \text{E}[e'](\rho[\text{if } \text{E}[e] \rho \in \{ K \} \times V \text{ then } \\
&\quad \text{snd}(\text{E}[e] \rho) \text{ else } \bot/x])
\end{align*}
\]

Semantics of types

Following [MPS86], we identify types with weak ideals over the semantic domain \( V \). A type environment \( \psi \in TEnv \) is a partial mapping from type variables to ideals and from Skolem type constructors to functions between ideals. The semantic interpretation of types, \( T : TExp \rightarrow TEnv \rightarrow \mathbb{S}(V) \), is defined as follows.
The universal and existential quantifications range over the set $\mathcal{R} \subseteq \mathcal{I}(V)$ of all ideals that do not contain wrong. Note that the sum in the definition of recursive types is actually a union, since the constructor tags are assumed to be distinct. It should also be noted that our interpretation does not handle ML’s nonregular, mutually recursive datatypes; it appears that the PER model described in [BM92] would provide an adequate interpretation.

Theorem 4 The semantic function for types is well-defined.

Proof: As in [MPS86]. We observe that $\lambda \eta \in \mathcal{I}(V). \sum \{ K_i \} \times T[\eta_i] (\psi[I/\beta])$ is always contractive, since cartesian product and sum of ideals are contractive; therefore, the fixed point of such a function exists.

Lemma 5 Let $\psi$ be a type environment such that for every $\alpha \in \text{Dom } \psi$, wrong $\in \psi(a)$. Then for every type scheme $\sigma$, wrong $\in T[\sigma \psi]$.

Proof: By structural induction on $\sigma$.

Lemma 6 [Substitution] $T[\sigma (\sigma' / \alpha)] = T[\sigma] (\psi[T[\sigma'] \psi / \alpha])$.

Proof: Again, by structural induction on $\sigma$.

Definition 1 [Semantic type judgment] Let $A$ be an assumption set, $e$ an expression, and $\sigma$ a type scheme. We define $\vdash_{\rho, \psi} A$ as meaning that $\text{Dom } A \subseteq \text{Dom } \rho$ and for every $x \in \text{Dom } A$, $\rho(x) \in T[A(x)] \psi$; further, we say $A \vdash_{\rho, \psi} e : \sigma$ iff $\vdash_{\rho, \psi} A$ implies $E[e] \rho \in T[\sigma] \psi$; and finally, $A \vdash e : \sigma$ means that for all $\rho \in \text{Env}$ and $\psi \in T\text{Env}$ we have $A \vdash_{\rho, \psi} e : \sigma$.

Theorem 7 [Semantic Soundness] If $A \vdash e : \tau$ then $A \vdash e : \tau$.

Proof: By induction on the size of the proof tree for $A \vdash e : \tau$. We need to consider each of the cases given by the type inference rules. Applying the inductive assumption and the typing judgments from the preceding
steps in the type derivation, we use the semantics of the types of the partial results of the evaluation. In each
of the cases below, choose \( \psi \) and \( \rho \) arbitrarily, such that \( \models_{\rho, \psi} A \). We include only the nonstandard cases.

Lemma 6 will be used with frequency.

\[ A \models \text{data} \ \forall \alpha_1 \ldots \alpha_n. \mu \beta. K_1 \eta_1 + \ldots + K_m \eta_m \ \text{in} \ e : \tau \]

The premise in the type derivation is \( A \models \sigma/K_1, \ldots, \sigma/K_m \models e : \tau \), where
\( \sigma = \forall \alpha_1 \ldots \alpha_n. \mu \beta. K_1 \eta_1 + \ldots + K_m \eta_m \). Since by definition, \( \models_{\rho, \psi} A \models \sigma/K_1, \ldots, \sigma/K_m \), we can
use the inductive assumption to obtain \( E[\text{data} \ \forall \alpha_1 \ldots \alpha_n. \chi \ \text{in} \ e \] \( \rho = E[e] \rho \in T[\tau] \psi \).

\[ A \models K : \tau \rightarrow \mu \beta. \Sigma[K\eta] \]

The last premise in the type derivation is \( \eta \ (\mu \beta. \Sigma[K\eta]) / \beta \leq \tau \), where \( \eta = \exists \gamma_1 \ldots \gamma_n \). By definition of instantiation of existential types, \( \tau = \tau (\gamma_j/\gamma_j, \mu \beta. \Sigma[K\eta]) / \beta \) for some types \( \tau_1, \ldots, \tau_n \).

First, choose an arbitrary \( v \in T[\tau] \psi \) and a finite \( a \leq v \). Now,

\[ a \in (T[\tau (\gamma_j/\gamma_j, \mu \beta. \Sigma[K\eta]) / \beta] \psi) \circ \]

\[ = (T[\tau (\mu \beta. \Sigma[K\eta]) / \beta] (\psi[T[\tau] \psi / \gamma_i])) \circ \]

\[ \subseteq \bigcup_{J_1, \ldots, J_n \in \mathbb{R}} (T[\tau (\mu \beta. \Sigma[K\eta]) / \beta] (\psi[J_j/\gamma_i])) \circ \]

\[ = (\bigcup_{J_1, \ldots, J_n \in \mathbb{R}} T[\tau (\mu \beta. \Sigma[K\eta]) / \beta] (\psi[J_j/\gamma_i])) \circ \]

\[ = (T[\eta (\mu \beta. \Sigma[K\eta]) / \beta] \psi) \circ . \]

Hence, \( v = \bigcup \{a \mid a \in T[\eta (\mu \beta. \Sigma[K\eta]) / \beta] \psi \), by closure of ideals under limits. Consequently,

\[ \langle K, v \rangle \in \{K\} \times T[\eta (\mu \beta. \Sigma[K\eta]) / \beta] \psi \]

\[ \subseteq \ldots + \{K\} \times T[\eta (\mu \beta. \Sigma[K\eta]) / \beta] \psi + \]

\[ = \ldots + \{K\} \times T[\eta] (\psi[T[\mu \beta. \Sigma[K\eta]] \psi / \beta]) + \]

\[ = T[\mu \beta. \Sigma[K\eta]] \psi . \]

Hence \( E[\langle K \rangle] \rho \in T[\tau \rightarrow \mu \beta. \Sigma[K\eta]] \psi \).

\[ A \models \text{is} \ K : (\mu \beta. \Sigma[K\eta]) \rightarrow \text{bool} \]

Choose an arbitrary \( v \in T[\mu \beta. \Sigma[K\eta]] \psi \). Clearly, \( (E[\text{is} \ K] \rho) v \in B \), whence
\( E[\text{is} \ K] \rho \in T[\mu \beta. \Sigma[K\eta]] \rightarrow \text{bool} \psi \).

\[ A \models \text{let} \ K \ x = e \ \text{in} \ e' : \tau' \]

We follow the proof in [MPS86]. The first premise in the type derivation is \( A \models e : \tau \), where
\( \tau = \mu \beta. \Sigma[K\eta] \) and \( \eta = \exists \gamma_1 \ldots \gamma_n \). Let \( \{\alpha_1, \ldots, \alpha_k\} = \text{FV}(\tau) \setminus \text{FV}(A) \). Then, for every
\( I_1, \ldots, I_k \in \mathbb{S} (V) \), \( \models_{\rho, \psi[I_j/\alpha_j]} A \) holds, since none of the \( \alpha_i \)'s are free in \( A \).

Let \( v = E[e] \rho \); by the inductive assumption, \( v \in T[\tau] (\psi[I_j/\alpha_j]) \). Consequently,

\[ \forall \in \bigcap_{I_1, \ldots, I_k \in \mathbb{R}} T[\tau] (\psi[I_j/\alpha_j]) \]

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First, consider the case \( \text{fst} (v) \neq K \). Then, by definition, \( E \llbracket \text{let } K \ x = e \ \text{in } e' \rrbracket \rho = \bot \), and we are done, since \( \bot \in T \llbracket \tau' \rrbracket \psi \).

In the more interesting, second case, \( \text{fst} (v) = K \). Then

\[
\text{snd} (v) \in \bigcap_{i_1, \ldots, i_k \in R} T \llbracket \tau' \rrbracket \left( \psi [l_i / \alpha_i, J_j / \gamma_j, T \llbracket \tau \rrbracket (\psi [l_i / \alpha_i]) / \beta] \right) + \ldots
\]

Let \( \alpha_1, \ldots, \alpha_h, h \leq k \), be those variables among \( \alpha_1, \ldots, \alpha_k \) that are free in \( \tau' [\tau / \beta] \).

We now choose a finite \( a \) such that \( a \leq \text{snd} (v) \), thus

\[
a \in \bigcap_{i_1, \ldots, i_k \in R} T \llbracket \tau' [\tau / \beta] \rrbracket \psi [l_i / \alpha_i, J_j / \gamma_j] \phi.
\]

By definition of set union and intersection, there exist functions \( f_1, \ldots, f_n \in \mathbb{S} (V)^h \rightarrow \mathbb{S} (V) \), such that

\[
a \in \bigcap_{i_1, \ldots, i_k \in R} T \llbracket \tau' [\tau / \beta] \rrbracket \psi [l_i / \alpha_i, f_j (l_i, \ldots, l_h) / \gamma_j] \phi.
\]

assuming that the \( \kappa_j \)'s are the ones generated by \( \text{skolem} (A, \eta [\tau / \beta]) \).

Since by definition of \( \text{skolem} \), none of the \( \kappa_j \)'s are free in \( A \), \( \models_{\rho, \psi [f_j / \kappa_j]} A \) holds and we can extend \( A \) and \( \rho \), obtaining \( \models_{\rho [a / x], \psi [f_j / \kappa_j]} A [\text{gen} (A, \text{skolem} (A, \eta [\tau / \beta])) / x] \).

We now apply the inductive assumption to the last premise,

\[
A [\text{gen} (A, \text{skolem} (A, \eta [\tau / \beta])) / x] \vdash e' : \tau',
\]

and obtain

\[
E \llbracket e' \rrbracket (\rho [a / x]) \in T \llbracket \tau' \rrbracket (\psi [f_j / \kappa_j]) = T \llbracket \tau' \rrbracket \psi,
\]

since \( FS (\tau') \subseteq FS (A) \). Finally,

\[
E \llbracket \text{let } K \ x = e \ \text{in } e' \rrbracket \rho = E \llbracket e' \rrbracket (\rho [\text{snd} (E \llbracket e \rrbracket \rho) / x])
\]

\[
= \bigcup \{ E \llbracket e' \rrbracket (\rho [a / x]) | a \text{ finite and } a \leq \text{snd} (E \llbracket e \rrbracket \rho) \},
\]

by the continuity of \( E \). The latter expression is in \( T \llbracket \tau' \rrbracket \psi \) by the closure of ideals under limits.

\[ \blacksquare \]

Corollary 8 [Semantic Soundness] If \( A \vdash e : \tau \), then \( E \llbracket e \rrbracket \rho \neq \text{wrong} \).

Proof: We apply Lemma 5 to the previous theorem.
References


Dynamic Typing in Polymorphic Languages

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1 Introduction

Dynamic types are sometimes used to palliate deficiencies in languages with static type systems. They can be used instead of polymorphic types, for example, to build heterogeneous lists; they are also exploited to simulate object-oriented techniques safely in languages that lack them, as when emulating methods with procedures. But dynamic types are of independent value, even when polymorphic types and objects are available. They provide a solution to a kind of computational incompleteness inherent to statically-typed languages, offering, for example, storage of persistent data, inter-process communication, type-dependent functions such as print, and the eval function.

Hence, there are situations in programming where one would like to use dynamic types even in the presence of arbitrarily advanced type features. In this paper we investigate the interplay of dynamic typing with polymorphism. Our study extends earlier work (see [1]) in allowing polymorphism, but keeps the same basic language constructs (dynamic and typecase) and the same style.

The interaction of polymorphism and dynamic types gives rise to problems in binding type variables. We find that these problems can be more clearly addressed in languages with explicit polymorphism. Even then, we encounter some perplexing difficulties (as indicated in [1]). In particular, there is no unique way to match the type tagging a dynamic value with a typecase pattern. Our solution consists in constraining the syntax of typecase patterns, providing static guarantees of unique solutions. The examples we have examined so far suggest that our restriction is not an impediment in practice. This solution applies also to languages with abstract data types, and it extends to languages with subtyping.

Drawing from the experience with explicit polymorphism, we consider languages with implicit polymorphism in the ML style. The same ideas can be used, with some interesting twists. In particular, we are led to introduce tuple variables, which stand for tuples of type variables.

In addition to [1], several recent studies have considered languages with dynamic types [14, 17, 23]. The work most relevant to ours is that of Leroy and Mauny, who define and investigate two extensions of ML with dynamic types. We compare their designs to ours in section 6.

Section 2 is a brief review of dynamic typing in simply-typed languages, based on [1]. Section 3 considers the general case of adding dynamic typing to a language with higher-order polymorphism [11]. An algorithmic formulation of the general framework is obtained in section 3.5 by restricting polymorphism to the second order and placing conditions on the patterns used in typecase expressions. Sections 4 and 5 discuss abstract data types and subtyping, respectively. Section 6 deals with a language with implicit polymorphism.

2 Review

The integration of static and dynamic typing is fairly straightforward for monomorphic languages. The simplest approach introduces a new base type Dynamic along with a dynamic expression for constructing values of type Dynamic and a typecase expression for inspecting them. The typechecking rules for these expressions are:

\[
\begin{align*}
\Gamma & \vdash a : T \\
\Gamma & \vdash \text{dynamic}(a : T) : \text{Dynamic} \\
\Gamma, x : P & \vdash b : T \\
\Gamma & \vdash e : T \\
\Gamma & \vdash \text{typecase} \ d \ of \ (x : P) \ b \ \text{else} \ e : T
\end{align*}
\]

(DYN-I)
(DYN-E)

The phrases \((x : P)\) and else are branch guards; \(P\) is a pattern—here, just a monomorphic type; \(b\) and \(e\) are branch bodies. For notational simplicity, we have
considered only typecase expressions with exactly one guarded branch and an else clause; typecase involving several patterns can be seen as syntactic sugar for several nested instances of the single-pattern typecase.

In the intended implementation, the compilation of a dynamic is a pair consisting of a value and its type:

\[
\text{compile}[\text{dynamic}(a:A)] = "<'\text{compile}[a]'\text{' grab}[a]">\]

The double quotes here indicate that the result of compile is a run-time structure; single quotes mark substructures to be built at compile time. The keyword "grab" indicates a metalevel shift: a compile-time data structure or routine is inserted into the run-time value. Because of this, Dynamic is particularly easy to implement in a bootstrapped compiler, where the run-time and compile-time structures coincide.

The typecase construct uses the compiler's type-match routine to compare the tag of a given dynamic to the branch guard:

\[
\text{compile}[\text{typecase } d \text{ of } (x:A) b \text{ else } e] = \\
\text{"let } d = '\text{compile}[d]' \text{ in} \\
\text{if } '\text{grab}[\text{typematch}](d) (\text{snd}(d)) \text{ then push}[x=fst(d)]; '\text{compile}[b]' \\
\text{else } '\text{compile}[e]'\"
\]

In languages with subtyping, it is common to use a subtype test in the typecase construct to give a less restrictive matching rule, allowing a tag to be a subtype of the guard type instead of requiring that the two match exactly.

Constructs analogous to dynamic and typecase have been provided in a number of languages, including Simula-67 [2], CLU [18], Cedar/Mesa [16], Amber [3], Modula-2+ [22], Oberon [25], and Modula-3 [9].

These constructs have surprising expressive power; for example, fixpoint operators can be defined at every type already in a simply typed lambda-calculus extended with Dynamic [1]. Important applications of dynamics include persistence and inter-address-space communication. For example, the following primitives provide input and output of a dynamic value from and to a stream:

\[
\text{extern } E \text{ writer } \text{ Dynamic } \rightarrow \text{ Unit} \\
\text{intern } E \text{ reader } \text{ Dynamic}
\]

Moreover, dynamics can be used to give a type for an eval primitive [12, 20]:

\[
\text{eval } E \text{ Exp } \rightarrow \text{ Dynamic}
\]

We obtain a much more expressive system by allowing typecase guards to contain pattern variables. For example, the following function takes two Dynamic arguments and attempts to apply the contents of the first (after checking that it is of functional type) to the contents of the second:

\[
\text{squarePolyFun } = \\
\lambda (df: \text{Dynamic}) \lambda (dg: \text{Dynamic}) \\
\text{typecase } df \text{ of } \\
\{ U, V \} (f: U \rightarrow V) \\
\lambda (W) \lambda (x: W) f[W](x) * f[W](x) \\
\text{else } \lambda (W) \lambda (x: W) 0
\]

3 Explicit Polymorphism

This formulation of dynamic types may be carried over almost unchanged to languages based on explicit polymorphism [11, 21]. For example, the following function checks that its argument df contains a polymorphic function \( f \) taking elements of any type into Int. It then creates a new polymorphic function that accepts a type and an argument of that type, instantiates \( f \) appropriately, applies it, and squares the result:

\[
\text{squarePolyFun } = \\
\lambda (df: \text{Dynamic}) \\
\text{typecase } df \text{ of } \\
\{ W \} (f: V(Z) \rightarrow \text{Int}) \\
\lambda (W) \lambda (x: W) f[W](x) * f[W](x) \\
\text{else } \lambda (W) \lambda (x: W) 0
\]

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Here the type abstraction operator is written $\lambda$. Type application is written with square brackets. The types of polymorphic functions begin with $\forall$. For example, $\forall(T)T \rightarrow T$ is the type of the polymorphic identity function, $\lambda(T) \lambda(x:T) x$.

### 3.1 Higher-order pattern variables

First-order pattern variables, by themselves, do not give us sufficient expressive power in matching against polymorphic types. For example, we might like to generalize the dynamic application example from section 2 so that it can accept a polymorphic function and instantiate it appropriately before applying it:

$$\text{dynApply2} = \lambda(df:\text{Dynamic}) \lambda(da:\text{Dynamic})$$

$$\text{typecase df of}$$

$$\begin{cases} 
\{ (f:\forall(Z)\rightarrow ?) \\
\text{typecase da of} \\
\{ W \ (a:W) \}
\end{cases}$$

$$\begin{cases} 
\text{dynamic}(f[W](a):?) \\
\text{else...} \\
\text{else dynApply(df)(da)} 
\end{cases}$$

But there is no single expression we can fill in for the domain of $f$ that will make $\text{dynApply2}$ apply to both:

$$\text{df} = \text{dynamic}(\lambda(Z) \lambda(x:Z \times Z) <\text{snd}(x),\text{fst}(x)>: \ldots)$$

$$\text{da} = \text{dynamic}(<3,4>: \ldots)$$

and:

$$\text{df} = \text{dynamic}((\lambda(Z) \lambda(x:Z \rightarrow Z) x): \ldots)$$

$$\text{da} = \text{dynamic}((\lambda(x:\text{Int}) x): \ldots)$$

Thus we are led to introducing higher-order pattern variables, which range over "pattern contexts"—patterns abstracted with respect to some collection of type variables. These suffice to express polymorphic dynamic application:

$$\text{dynApply2} = \lambda(df:\text{Dynamic}) \lambda(da:\text{Dynamic})$$

$$\begin{cases} 
\text{typecase df of} \\
\{ F,G \} (f:\forall(Z)F(Z) \rightarrow G(Z)) \\
\text{typecase da of} \\
\{ W \ (a:F(W)) \}
\end{cases}$$

$$\begin{cases} 
\text{dynamic}(f[W](a):G(W)) \\
\text{else...} \\
\text{else dynApply(df)(da)} 
\end{cases}$$

For example, if:

$$\text{df} = \text{dynamic}(\text{id}:\forall(A)A \rightarrow A)$$

$$\text{da} = \text{dynamic}(3: \text{Int})$$

then the typecase expressions match as follows:

<table>
<thead>
<tr>
<th>Tag</th>
<th>Pattern</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\forall(A)A \rightarrow A$</td>
<td>$\forall(Z)F(Z) \rightarrow G(Z)$</td>
<td>${ F = \Lambda(x)I, G = \Lambda(x)I }$</td>
</tr>
</tbody>
</table>

and the result of the application $\text{dynApply2}(df)(da)$ is $\text{dynamic}((\text{id}[\text{Int}](3): \text{Int})$.

### 3.2 Syntax

We now formalize dynamic types within the context of a higher-order polymorphic $\lambda$-calculus, $F_\omega$ [11]. The syntax of $F_\omega$ with type $\text{Dynamic}$ is given in Figure 1.

In examples we also use base types, cartesian products, and labeled records in types and patterns, but we omit these in the formal treatment.

We regard as identical any pair of formulas that differ only in the names of bound variables. For brevity, we sometimes omit kinding declarations of the form "Type" and empty pattern-variable bindings. Also, it is technically convenient to write the pattern variables bound by a typecase expression as a syntactic part of the pattern, rather than putting them in front of the guard as we have done in the examples. Thus, $\text{typecase e1 of } \{ W \}(x:T)e2$ else $e3$ should be read formally as $\text{typecase e1 of } (x:\forall(T)T)e2$ else $e3$.

### 3.3 Tag Closure

One critical design decision for a programming language with type $\text{Dynamic}$ is the question of whether type tags must be closed (except for occurrences of pattern variables), or whether they may mention universally bound type variables from the surrounding context.

In the simplest scenario, $\text{dynamic}(a:A)$ is legal only when $A$ is a closed (but possibly polymorphic) type. Similarly, we would require that the guard in a typecase expression be a closed type.

If the closure restriction is not instituted, then types must actually be passed as arguments to polymorphic functions at run time, so that code can be compiled for expressions like:

$$\lambda(x) \lambda(x:Z) \text{dynamic}(<x,x>:Z \times Z)$$

where the type $Z \times Z$ must be generated at run time. For languages such as ML, where type information is not retained at run time, the closure restriction becomes essential (see section 6). For now, we consider the general case where tags may contain free type variables.

### 3.4 Definiteness

The most simple-minded formulation of higher-order pattern variables may seem to provide adequate expressive power, but it is not sufficiently constrained to lead to a viable language design. The problem is that there is no guarantee of unique matches of patterns against tags. For example, when the pattern $F(\text{Int})$ is matched
Figure 1: Syntax of $F_\omega$ terms

```plaintext
K ::= Type
    | K → K

T ::= A
    | T → T
    | \forall (A : K)T
    | \Lambda (A : K)T
    | T(T)
    | Dynamic

P ::= \{V_1 : K_1, \ldots, V_n : K_n\} T

e ::= z
    | \lambda (z : T) e
    | e(e)
    | \lambda (A : K) e
    | e[T]
    | dynamic(e : T)
    | typecase e of (z : P) e else e
```

against the tag Bool, the pattern variable $F$ is forced to be $\Lambda (X)\text{Bool}$. But when the same pattern is matched against the tag Int, we find that $F$ can be either $\Lambda (X)X$ or $\Lambda (X)\text{Int}$. There is no reasonable way to choose.

Worse yet, consider $F(W)$ or $F(W \rightarrow \text{Int})$ for a pattern variable $W$.

These problems compel us to introduce restrictions on the form of patterns. We may require that a pattern matches a tag in at most one way at run time, and fail otherwise. But this leads to unpredictable matching failures. Therefore we prefer a stronger condition.

Informally, we want to allow only patterns that match each tag in at most one way. This condition is called definiteness. For example, assume that the type variables $A$ and $B$ and the operator variable $H$ are bound in the current context, and consider the following patterns:

Valid: $\{V\}$ $H(V)$

since, at run time, if $V$ appears in the expression to which $H$ is bound, then matching is the usual for first-order pattern variables; and otherwise matching leaves $V$ completely undetermined, and we set it to a new type constant. Patterns of this form are used in many of our examples, so we want to consider them definite even though they may sometimes leave $V$ unconstrained.

Invalid: $\{F\}$ $F(A)$

because, for example, it can match the tag $A \rightarrow A$ in four ways, instantiating $F$ to any of: $\Lambda (C)C \rightarrow C$, $\Lambda (C)C \rightarrow A$, $\Lambda (C)A \rightarrow C$, $\Lambda (C)A \rightarrow A$.

Valid: $\{F\}$ $\forall (X)F(X)$

since, when the scope of $X$ is narrower than that of $F$, we can only match the tag $\forall (Y)Y \rightarrow Y$ by instantiating $F$ to $\Lambda (C)C \rightarrow C$.

Valid: $\{F\}$ $(\forall (X)F(X)) \rightarrow F(A)$

since the first occurrence of $F$ determines its value.

Valid: $\{F, V\}$ $(\forall (X)F(X)) \rightarrow F(V)$

since $F$ can be matched first and then considered "already bound" at the defining occurrence of $V$.

Invalid: $\{F, G\}$ $\forall (X)F(G(X)) \rightarrow G(F(X))$

since neither $F$ nor $G$ can be considered "already bound" unless the other is bound first.

Note that definiteness of patterns cannot be checked locally. For example,

Valid: $\{F\}$ $F(\text{Int}) \rightarrow F(\text{Bool})$

is definite, although neither occurrence of $F$ would be definite in isolation.

The definiteness condition can be formalized, and then one can put a definiteness requirement in the type-checking rule for typecase, so that only programs with definite patterns are legal.

Unfortunately, the notion of definiteness does not suggest a typechecking algorithm in any straightforward way. A related problem is that we have no algorithm for the run-time operation of matching patterns against tags. Indeed, it is not known whether the general case of higher-order matching is decidable. Even the second- and third-order cases, whose decidability has been established [15, 7, 8], lead to algorithms too inefficient to be of practical use in implementing typecase.
3.5 Second-order polymorphism

To obtain a practical language design, we need a restriction of our general treatment for which efficient typechecking and matching algorithms can be given. We begin by considering the fragment of $F_w$ with only second-order polymorphism. This restriction is mostly a matter of convenience, and it seems possible that the approach described below applies to the full $F_w$.

The syntax of System $F$ (the second-order polymorphic lambda-calculus) with Dynamic is given by the following restriction of $F_w$ with Dynamic. We show only the lines that differ.

$$K ::= \ldots$$
$$| \text{Type} \rightarrow K \quad \text{type operators}$$

$$T ::= \ldots$$
$$| \forall(A)T \quad \text{quantified types}$$
$$| \Lambda(A)T \quad \text{type operators}$$
$$| \ldots$$

$$P ::= \ldots$$

$$e ::= \ldots$$
$$| \lambda(A)e \quad \text{type abstraction}$$
$$| \ldots$$

Here, kinds other than Type are used only to specify the functionality of pattern variables; abstractions and applications at the level of types are used only to describe patterns. Since every kind has the form

$$\text{Type} \rightarrow \ldots \rightarrow \text{Type},$$

we can simply say that a pattern variable with this kind has arity $n$.

A pattern $\{V_1 : K_1, \ldots, V_n : K_n\} T$ is ordered if there is some total ordering $<$ of the pattern variables $V_1, \ldots, V_n$ such that each $V_i$ has a defining occurrence, that is, a subterm occurrence $U \equiv V_i \cdot A_1 \cdots A_p$ in $T$ such that:

1. $U$ does not appear in an argument to a pattern variable $V_j$ where $V_j \geq V_i$ (i.e., there is no occurrence $U' \equiv V_j \cdot Q$ with $U$ a proper subphrase of $U''$ and $V_j \geq V_i$);
2. $V_i$ is fully applied (i.e., the arity of $V_i$ is $p$);
3. the $A_j$'s are pairwise distinct; and
4. all of the $A_j$'s have narrower scope than $V_i$ (i.e., $A_j \notin \text{FV}(T)$).

Note that this condition can be checked statically.

*Ordered matching* is a matching algorithm for ordered patterns; given an ordered pattern, this algorithm instantiates variables according to one of the orders that make the pattern ordered. We believe that ordered patterns are always definite, and that ordered matching is correct, that is, it terminates on every input and always yields the same solution independently of the order of variable instantiations. Hence we replace the definiteness condition with the ordered condition in typechecking, and in evaluation we adopt ordered matching. We omit a detailed description of ordered matching; section 6.3 contains a similar algorithm in a somewhat different context.

4 Abstract Types

The interaction between the use of Dynamic and abstract data types gives rise to a puzzling design issue: should the type tag of a dynamically typed value containing an element of an abstract type be matched abstractly or concretely? There are good arguments for both choices:

- Abstract matching protects the identity of "hidden" representation types and prevents accidental matches in cases where several abstract types happen to have the same representation.

- Transparent matching allows a more permissive style of programming, where a dynamically typed value of some abstract type is considered to be a value of a different version of "the same" abstract type. This flexibility is critical in many situations. For example, a program may create disk files containing dynamic values, which should remain usable even after the program is recompiled, or two programs on different machines may want to exchange abstract data in the form of dynamically typed values.

By viewing abstract types formally as existential types [19], we can see exactly where the difference between these two solutions lies and suggest a generalization of existential types that supports both. (Existential types can in turn be coded using universal types, with this coding, our design for dynamic types of the previous sections yields the second solution.)

To add existential types to the variant of $F_w$ defined in the previous section, we extend the syntax of types and terms as in Figure 2.

The typechecking rules for pack and open are:

$$\frac{S \equiv_b \exists(A : K) \quad \Gamma \vdash e \in [R/A]T}{\Gamma \vdash \text{pack } e \text{ as } S \text{ hiding } R \in S} \quad \text{(Pack)}$$

$$\frac{\Gamma \vdash e \in \exists(A : K) \quad \Gamma, A : K, x : S \vdash b \in T}{\Gamma \vdash \text{open } e \text{ as } [A, x] \text{ in } b \in T} \quad \text{(Open)}$$

A typical example where an element of an abstract type is packed into a Dynamic is:

96
T ::= ... | \exists (A : K) T  \quad \text{existential types}

e ::= ... | \text{pack } e \text{ as } T \text{ hiding } T | \text{open } e \text{ as } [A, z] \text{ in } e \quad \text{packing (existential introduction)}
| \text{unpacking (existential elimination)}

Figure 2: Extended syntax with existential types

let stackpack =
  pack = λ(s: IntList)
       \lambda(i: Int) \text{ cons}(i)(s),
  pop = \lambda(s: IntList) \text{ cdr}(s),
  top = \lambda(s: IntList) \text{ car}(s),
  new = nil
as Some(X)

in
  open stackpack as [Stack, stackops] in
  let dstack =
    dynamic
    (stackops.push(stackops.new)(5): Stack)
  in
  typecase dstack of
    (s: Stack) stackops.top(s)
    else ... 0

Note that this sort of example depends critically on the use of open type tags. As above, open tags must be implemented using run-time types. The evaluation of pack must construct a value that carries the representation type.

We have a choice in the evaluation rule for the open expression:

- We can evaluate the expression open e as [R, z] in b by replacing the representation type variable R by the actual representation type obtained by evaluating e.
- Alternatively, we can replace R by a new type constant.

Without Dynamic, the difference between these rules cannot be detected. But with Dynamic we get different behaviors. Since both behaviors are desirable, we may choose to introduce an extended open form that provides separate names for the abstract and transparent versions of the representation type:

\[
\Gamma \vdash e \in \exists (H : K) S \quad H \not\in \text{FV}(T) \\
\Gamma, H : K, z : S \vdash [H/R]b \in T \\
\Gamma \vdash (\text{open } e \text{ as } [R, H, z] \text{ in } b) \in T \quad \text{(OPEN)}
\]

In the body of b, we can build dynamic values with tags R or H; a typecase on the former could investigate the representation type while a typecase on the latter could not violate the type abstraction.

5 Subtyping

In simple languages with subtyping (e.g., [3, 9]) it is natural to extend typecase to perform a subtyping test instead of an exact match. Consider for example:

let dx = dynamic(3:Nat)

in
  typecase dx of
    \{} (x: Int) ...
    else ... 0

The first typecase branch is taken: although the tag of dx, Nat, is different from Int, we have Nat \leq Int.

Unfortunately, this idea runs into difficulties when applied to more complex languages. In general, there does not exist a most general instantiation for pattern variables when a subtype-match is performed. For example, consider the pattern V → V and the problem of subtype-matching (Int→Nat) \leq (V → V). Both Int→Int and Nat→Nat are instances of V→V and supertypes of Int→Nat, but they are incomparable. Even when the pattern is covariant there may be no most general match. Given a pattern V × V, there may be a type A × B such that A and B have no least upper bound, and so there may be no best instantiation for V. This can happen, for example, in a system with bounded quantifiers [6, 10], and in systems where the collection of base types does not form an upper semi-lattice. Linear patterns (where each pattern variable occurs at most once) avoid these problems, but we find linearity too restrictive.

Therefore, we take a different approach that works in general and fits well with the language described in section 3.2. We intend to extend System F with subtyping along the lines of [5]. In order to incorporate also the higher-order pattern variables, we resort to powerkinds [4]. The kind structure of section 3.2 is therefore extended as follows:

\[
K ::= \text{Type} \\
| K \rightarrow K \\
| \text{Power}(K)(T) \quad \text{(where } T : K)\]

Informally, \text{Power}(Type)(T) is the collection of all subtypes of T, and \text{Power}(K→K')(F) is the collection of
all operators of kind $K \rightarrow K'$ that are pointwise in subtype relation with $F$. Subtyping ($\leq$) is introduced by interpreting:

$A \leq B : K$ as $A : \text{Power}(K)(B)$, where $A, B : K$

$F \leq G : (K \rightarrow K')$ as $F(X) \leq G(X) : K'$, for all $X : K$, where $F, G : (K \rightarrow K')$

The axiomatization of $\text{Power}(K)(T)$ [4] is designed to induce the expected subtyping rules. For example, $A : \text{Power}(A)$ says that $A \leq A$.

As in section 3.5, we limit kinds to appear only in patterns, although we may allow bounded quantifiers $\forall(X \leq T)T'$ since they can be handled easily. Because of power-kinds, we can now write patterns such as:

```plaintext
typecase dx of
  \{V, W \leq (V \times V)\} (x : W) ...
  (that is: \{V : Type, W : Power(Type)(V \times V)\} (x : W))
  else ...
```

Each branch guard is used in typechecking the corresponding branch body. The shape of branch guards is $\{V_1 : P_1, \ldots, V_n : P_n\}(x : P)$ where each $V_i$ may occur in the $P_j$ with $j > i$ and in $P$. This shape fits within the normal format of typing environments, and hence introduces no difficulties for static typechecking.

Next we consider the dynamic semantics of `typecase` in presence of subtyping. The idea is to preserve the previous notion that `typecase` performs exact type matches at run time. Subtyping is introduced as a sequence of additional constraints to be checked at run time only after matching. These constraints are easily checked because, by the time they are evaluated, all the pattern variables have been fully instantiated (perhaps to undetermined types, as discussed in section 3). In the example above, suppose that the tag of $dx$ is $(\text{Nat} \times \text{Int}) \times \text{Int}$; then we have the instantiations $W = \text{Nat} \times \text{Int}$ and $V = \text{Int}$. When the matching is completed, we successfully check that $W \leq (V \times V)$.

Some examples will illustrate the additional flexibility obtained with subtyping. First we show how to emulate simple monomorphic languages with subtyping but without pattern variables, where `typecase` performs a subtype test. The first example of this section can be reformulated as:

```plaintext
typecase dx of
  \{V \leq \text{Int}\}(x : V) f[V](x)
  else ...
```

where $f : \forall(V \leq \text{Int})W \rightarrow \text{Int}$. In this example, the tag of $dx$ can be any subtype of $\text{Int}$. Note that the assumption $V \leq \text{Int}$ is used statically to typecheck $f[V](x)$.

The next example is similar to `dynApply` in section 2, but the type of the argument can be any subtype of the domain of the function:

```plaintext
\text{typecase df of}
  \{V, W\} (f : V \rightarrow W)
  \text{typecase da of}
    \{V' \leq V\} (a : V')
      \text{dynamic}(f(a) : W)
    else ...
  else ...
```

With polymorphic tag types, or with polymorphic pattern types with only first-order pattern variables, nothing new happens except that the matching and subtype tests must be the adequate ones for polymorphism.

The next degree of complexity is introduced by higher-order pattern variables. Just as we had $V' \leq V$, a subtype constraint between two first-order pattern variables, we may have $F \leq G : (K \rightarrow K')$ for two higher-order pattern variables $F, G : (K \rightarrow K')$. As mentioned above, the inclusion is intended pointwise: $F \leq G$ iff $F(X) \leq G(X) : K'$ under the assumption $X : K$.

Another form of dynamic application provides an example of higher-order matches with subtyping:

```plaintext
\text{typecase df of}
  \{F, G : \forall(X)G(X) \rightarrow H(X)\}
  \text{typecase dg of}
    \{F : \forall(Y)F(Y) \rightarrow \text{G'}(Y)\}
      \text{dynamic}
        ((\lambda(Z) f[Z] \text{og}[Z]) : \forall(Z)F(Z) \rightarrow H(Z))
    else ...
  else ...
```

Finally, dynamic composition calls for a constraint of the form $G' \leq G$:

```plaintext
\text{typecase df of}
  \{G, H : \forall(X)G(X) \rightarrow H(X)\}
  \text{typecase dg of}
    \{F : \forall(Y)F(Y) \rightarrow \text{G'}(Y)\}
      \text{dynamic}
        ((\lambda(Z) f[Z] \text{og}[Z]) : \forall(Z)F(Z) \rightarrow H(Z))
    else ...
  else ...
```

This example generalizes to functions of bounded polymorphic types, such as $\forall(X \leq A)G(X) \rightarrow H(X)$.

## 6 Implicit Polymorphism

In this section we investigate dynamics in an implicitly typed language, namely ML. First we show that the general treatment of dynamics for explicitly typed languages directly applies to ML, providing explicitly tagged dynamics in an untyped language. This solution is not in the spirit of ML, and all the rest of the section will be devoted to the study of implicitly tagged dynamics in ML.

In the obvious extension of ML, types can still be inferred for all constructs but dynamics; the user needs
to provide type information when creating or reading dynamics. For instance, let us consider the program:

\[
\text{twice} = \text{dynamic} \quad (\lambda(f) \lambda(x) \, f(f \, x) : \forall(A)(A\rightarrow A) \rightarrow (A\rightarrow A))
\]

First, the type scheme \(\forall(A)(A\rightarrow A) \rightarrow (A\rightarrow A)\) is inferred for \(\lambda(f) \lambda(x) \, f(f \, x)\) as if it were to be let-bound. Then we check that this type scheme has no free variables and is more general than the required tag \(\forall(A)(A\rightarrow A) \rightarrow (A\rightarrow A)\). Conversely, when the extraction of a value from a dynamic succeeds, it is given the type scheme of its tag as if it had been let-bound. Thus, all instances of the value can be used with different instances of the tag as in

\[
\text{foo} = \lambda(df) \quad \text{typecase df of} \quad (f: \forall(A)(A\rightarrow A) \rightarrow (A\rightarrow A)) \quad <f \text{ succ, } f \text{ not}> \quad \text{else} \ldots
\]

where succ and not are the successor function on integers and the negation function on booleans.

This works perfectly. However, it requires explicit type information in dynamic patterns, which is not in the spirit of ML. Since the ML typechecker can infer most general types for expressions, one would expect the compiler to tag dynamic values with their principal types. For instance, the user writes

\[
\text{twice} = \text{dynamic} (\lambda(f) \lambda(x) f(f \, x))
\]

and the dynamic is tagged with \(\forall(A)(A\rightarrow A) \rightarrow (A\rightarrow A)\).

However, there is a difficulty with the program

\[
\text{apply} = \text{dynamic} (\lambda(f) \lambda(x) f \, x)
\]

Should the dynamic’s tag be \(\forall(A,B)(A\rightarrow B) \rightarrow (A\rightarrow B)\) or \(\forall(B,A)(A\rightarrow B) \rightarrow (A\rightarrow B)\)? As typecase is defined, it succeeds if the tag exactly matches the pattern, including quantifiers. With implicit tagging, the order of quantifiers should not matter.

Moreover, since the tag of \text{twice} is more general than the pattern of the function \text{foo}, an ML programmer would probably expect that \text{twice} can be passed to \text{foo}. This is also justified by the fact that the typechecker could have built a dynamic with a weaker tag, and the typecase would have succeeded. That is, in ML, the typecase would be expected to succeed if an instance of the tag matches the pattern. This principle is called tag instantiation. Dynamics with tag instantiation but no pattern variables have been implemented in the language CAML [24]. The dynamics studied by Leroy and Mauny [17] have tag instantiation and first-order pattern variables. First-order pattern variables are not powerful enough to type some reasonable examples, such as the \text{applyTwice} function shown later. Below we describe a version of dynamics for ML with tag instantiation and second-order pattern variables.

### 6.1 Tuple variables

Tag instantiation and second-order pattern variables do not fit well together. The difficulty comes from the merging of two features:

- As in the pattern \{F\} (f: \forall(A)F(A) \rightarrow A), second-order pattern variables may depend on universal variables.

- Tag instantiation requires that if a typecase succeeds, then it also succeeds for a dynamic with an argument that has a more general tag. The tag \(\forall(A)(A\times A) \rightarrow A\) matches the previous pattern. So should the tag \(\forall(A,B)(A\times B) \rightarrow A\). But F is not supposed to depend on B!

Because of tag instantiation, polymorphic pattern variables may always depend on more variables than the ones explicitly mentioned. We capture all variables that appear in the tag but that do not correspond to variables in the pattern into a tuple of variables P. The dependence of pattern variable F on all universal variables is written F(P), even though the exact set of variables in P is not statically known. The tuple variable P will be dynamically instantiated to the tuple of all variables of the tag not matched with variables of the pattern. In particular, if the pattern is F(P; A), then P will never contain A.

For instance, the pattern \{F\} (f: \forall(A)F(A) \rightarrow A) should be written \{F\} (f: \forall(A)F(P; A) \rightarrow A), so that tag instantiation is possible. The tag \(\forall(A)(A\times A) \rightarrow A\) matches this pattern for an empty tuple. The tag \(\forall(A,B)(A\times B) \rightarrow A\) matches it for a one-element tuple, namely (B).

Tuple variables bound in different patterns may be instantiated to tuples with different numbers of variables, as in the example just given. Because of such size considerations, it is not always possible to use a tuple variable as argument to an operator, since it may expect an argument of different size. We use tuple sorts in order to guarantee that type expressions are well formed. Formally, a typecase with explicit information should be written, for instance:

\[
\{p \text{ Tuple}, F:p\rightarrow \text{Type}\rightarrow \text{Type}\} \\
(f: \forall(P,p,A:\text{Type})F(P; A) \rightarrow A)
\]

where \(F(A_0; A_1, \ldots, A_n)\) stands for a fully applied pattern variable \(F(A_0)(A_1) \ldots (A_n)\); this notation reminds that the first argument \(A_0\) is a tuple. The sort variable \(p\) is to be bound at run time. However, it is not necessary to write the sorts in programs since a typechecker can easily infer them.

### 6.2 Description of the language

We assume given a denumerable collection of tuple sorts, written \(\pi, \pi', \ldots\), and a sort Type. Then the
sorts are:

\[ k ::= \pi \mid \text{Type} \]

atomic sorts

\[ K ::= k \mid k \to K \]

sorts

Types are:

\[ T ::= \text{Dynamic} \mid A \mid T(T) \mid T \to T \]

types

\[ P ::= \{A_1, \ldots, A_n\}T \]

patterns

\[ S ::= \forall(A_1 : K_1, \ldots, A_n : K_n) T \]

type schemes

In traditional ML style, we have left quantifiers implicit in types and hence in patterns. The formation of types, patterns, and contexts is controlled by judgements of the form:

\[ \Gamma \vdash T \in K \]

\[ \Gamma \vdash S \in K \]

\[ \Gamma \vdash P \in K \]

where contexts are:

\[ \Gamma ::= \emptyset \]

\[ \Gamma, \pi \]

tuple sort declaration

\[ \Gamma, F : K \]

symbol declaration

Formation rules ensure that type variables are always bound in the proper context, with a sort consistent with their uses. For instance, we have the rule:

\[ \frac{\Gamma \vdash T : k \quad \Gamma \vdash T' : k \to K}{\Gamma \vdash T'(T) : K} \quad \text{(SORT-APP)} \]

In examples only, we help the reader by using different letters for variables of different sorts: type variables of tuple sorts are written \( P \) and \( Q \) and type variables of sort Type are written \( A, B, \) etc. We also use \( F, G, \) and \( R \) for pattern variables.

Patterns are pairs of a set of pattern variables \( \{V_1, \ldots, V_n\} \) and a type \( T \). They are well formed if the signature of all \( V_i \)'s if of the form \( \pi \to K \) for the same tuple sort \( \pi \). The exact rule for pattern formation is:

\[ \frac{\pi, A_0, A_1, \ldots, A_n \notin \Gamma \quad \pi, V_1 : \pi \to K_1, \ldots, V_n : \pi \to K_n \quad A_0 : \pi, A_1 : \text{Type}, \ldots, A_n : \text{Type} \quad T \in \text{Type}}{\Gamma, \pi \vdash \{V_1 : \pi \to K_1, \ldots, V_n : \pi \to K_n\} T \in \forall(A_0 : \pi, A_1 : \text{Type}, \ldots, A_n : \text{Type}) T} \quad \text{(PATTERN-APP)} \]

In particular, there is exactly one pattern variable of each tuple sort per pattern.

Again, we would like to guarantee definiteness of patterns, and we impose the sufficient condition that patterns be ordered. Ordered patterns are those that satisfy the conditions given in section 3.5, and in addition all non-pattern free variables of sort Type must appear at least once outside of all pattern variables in the pattern. In the context of ML, our definiteness requirement is reminiscent of the type-explication restriction imposed on signatures in Standard ML (see section 7.7 of [13]). Pattern variables are bound at the beginning of the typecase, and their scope is the typecase in which they have been introduced. All other free variables are bound at the beginning of the pattern and their scope is the pattern.

Expressions are:

\[ e ::= x \mid \lambda(x) e \mid e e \]

\[ \text{dynamic}(e) \]

\[ \text{typecase } e \text{ of } \{V_1 : K_1, \ldots, V_n : K_n\} \quad (x : P) e \text{ else } e \]

Type inference Pattern variables behave as local type symbols in ML. Typechecking with local type symbols implies an extension of judgement contexts in order to control the scope of type symbols:

\[ \Gamma ::= \ldots \]

\[ \Gamma, \pi : S \]

variable type assignment

The typing judgements are \( \Gamma \vdash e : T \).

The "instance" rule of ML becomes:

\[ \frac{\Gamma \vdash T : \text{Type} \quad T \text{ is an instance of } S}{\Gamma \vdash e : T} \quad \text{(INST)} \]

It says that instances have to be well formed in the current context, which prevents us from using local symbols out of their scope.

Since we do not want to carry types at run time, we require that the types of values to become dynamic be closed, and then tags can be statically compiled.

\[ \frac{\Gamma \vdash e \in S \quad S \text{ is closed}}{\Gamma \vdash \text{dynamic}(e) \in \text{Dynamic}} \quad \text{(DYN-I)} \]

This rule may destroy the principal typing property of ML. If the principal type of an expression \( e \) is \( S \) and \( S \) is not closed, then typing \( \text{dynamic}(e) \) requires that free variables of \( S \) are instantiated by ground types. However, the set of closed instances of a principal type that is not closed does not have a principal element.

We want to avoid such situations, since the nonexistence of a principal type corresponds to an ambiguity concerning the tag that a dynamic value should carry. Therefore, we say that a program is not well typed if it has no principal typing derivation.

Type inference is realized by the same algorithm as in ML but delaying tag-closure checking to the end of typechecking (by gathering free variables of types of dynamic values in a list, for instance). If one of these variables is still free at the end of typechecking, then there exists no principal derivation, and the program is not well typed.
The rule for \texttt{typecase} is:

\[
\begin{align*}
\Gamma &\vdash d \in \text{Dynamic} & \Gamma &\vdash \epsilon' \in B \\
\Gamma, \pi \vdash \{V_1 : K_1, \ldots, V_n : K_n\} T \in S \\
\Gamma, x, V_1 : K_1, \ldots, V_n : K_n, x : S &\vdash e \in B \\
\Gamma, \mathcal{P} : \pi &\vdash \text{typecase } d \text{ of } \\
\{V_1, \ldots, V_n\} &\vdash (x : T) e \text{ else } \epsilon' \in B
\end{align*}
\]

The other rules of ML are unchanged.

6.3 Evaluation

Compilation is easily decomposed into two phases. The first phase translates ML into a variant, called ML\textsuperscript{-}, where dynamics are explicitly typed; this translation requires a bit of inference. ML\textsuperscript{-} differs form ML only in its \texttt{dynamic} construct:

\[
e := \ldots | \texttt{dynamic}(e : S)
\]

and its typing rule:

\[
\begin{align*}
\Gamma &\vdash a \in S & S \text{ is closed} \\
\Gamma &\vdash \texttt{dynamic}(a : S) \in \text{Dynamic}
\end{align*}
\]

(DYN-I)

The translation of an ML program \( e \) into ML\textsuperscript{-} is any ML\textsuperscript{-} program \( M \) whose principal type derivation is also a principal type derivation for \( e \). This defines \( M \) uniquely (types being equal up to alpha-conversion).

The type reconstruction algorithm is a trivial adaptation of the usual type inference algorithm. The semantics of an ML program \( e \) is the semantics of its translation into ML\textsuperscript{-}.

The evaluation rules are mostly standard. The only interesting one is for \texttt{typecase}, as it involves new methods for matching and pattern-variable instantiation.

Matching is not quite as usual, since it allows tag instantiation, and it also has to deal with tuple variables. Its inputs are a pattern \( \{V_1 : K_1, \ldots, V_n : K_n\} \) \( T \) and a tag, that is, a closed type \( \texttt{V}(\alpha_1, \ldots, \alpha_n) \cdot \tau \). The pattern variables are the \( V_i \)'s, and the universal variables are the remaining free variables of \( T \). The set of variables that occur in the tag (the \( \alpha_i \)'s) can increase during tag instantiation. The algorithm returns a substitution \( \mu \) with domain the pattern variables, such that there exists a substitution \( \mu' \) with domain the tag variables, and with \( \mu'(\tau) = \mu(T) \).

We describe the algorithm as transformations on sets of unification constraints called unificands; the transformations keep unchanged the set of substitutions that satisfy the constraints. The substitutions that we consider can instantiate both pattern and tag variables, but not universal variables.

The metavariable \( T \) still stands for any type, and \( \tau \) stands for a type that does not contain pattern variables. The atomic constraints are pairs, \( T = \tau \) or \( \tau = T \). The pairs \( T = \tau \) and \( \tau = T \) are considered equal. A substitution \( \mu \) is a solution of an atomic constraint if it unifies both sides. In addition, the constant \( \bot \) is used to represent failure; it is the atomic constraint with no solution.

In general, a unificand \( U \) is an atomic constraint, the conjunction of two unificands \( U' \land U'' \), or the existential unificand \( \exists a. U' \). The solution set of \( U' \land U'' \) is the intersection of the solution sets of \( U' \) and \( U'' \). The solution set of the existential unificand \( \exists a. U' \) is the set of solutions of \( U' \) restricted to variables distinct from \( a \). We identify unificands up to: commutativity and associativity of conjunction, renaming of variables bound by \( \exists \)'s, exchange of consecutive \( \exists \)'s, and removal of vacuous \( \exists \)'s.

Two unificands \( U \) and \( V \) are equivalent if they have the same set of solutions. This obviously defines an equivalence relation on unificands, and in fact a congruence.

We reduce the original matching problem to that of finding the solutions of the unificand \( \exists \text{FV}(\tau). (T \equiv \tau) \). In order to solve this problem, we now give a list of equivalences between unificands—the unificand on top is always equivalent to the one at the bottom. Tag variables are written \( \alpha \); \( C \) and \( C' \) are constant symbols, and always occur fully applied; and \( X \) is either a universal variable \( A \) or a pattern variable \( V \). The set of all variables is \( V \).

- **Decomposition**

\[
\begin{align*}
&\frac{C(T_i) \equiv C(\tau_i)}{\wedge (T_i \equiv \tau'_i)} & & \text{if } C \neq C' \\
&\frac{C(T_i) \equiv C'({\tau_j})}{\bot}
\end{align*}
\]

- **Instantiation**

\[
\exists a_i. (\alpha \equiv C(a_i) \land \wedge (T_i \equiv a_i))
\]

\[
\begin{align*}
V(A_0; a_1, \ldots, a_n) \equiv \tau \\
V \equiv \Lambda(A_0; a_1, \ldots, a_n). \tau
\end{align*}
\]

- **Propagation**

\[
\begin{align*}
X \equiv \tau \land M \\
X \equiv \tau \land [\tau / X]M
\end{align*}
\]

\[
\alpha \equiv \tau \\
\bot
\]

- **Universal-variable restriction**

\[
\begin{align*}
&\frac{A_i \equiv \alpha \land A_j \equiv \alpha}{\bot} & & \text{if } \alpha \notin \text{FV}(\tau), \tau \notin V \\
A \equiv \tau \\
\tau \notin V
\end{align*}
\]

- **Existential simplifications**

\[
\begin{align*}
&U \land (\exists a. U') \equiv \exists a. (U \land U') & & \alpha \notin \text{FV}(U) \\
&\exists a. (\alpha \equiv \tau \land U) \equiv \exists a. U & & \alpha \notin \text{FV}(\tau) \cup \text{FV}(U)
\end{align*}
\]

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• Trivial constraints

\[
\frac{\alpha \equiv \alpha \land U}{U} \quad \frac{\bot \land U}{\bot}
\]

These equivalences can be used as rewriting rules. All rules are oriented from top to bottom; one step of rewriting is the application of exactly one rule; applying the rules in any order always terminates. When successful, this process produces canonical unificands of the form:

\[
\exists \alpha_k \left( \bigwedge (A_i = \alpha_i) \land \bigwedge (V_j = \tau_j) \right)
\]

A unificand that cannot be reduced and that is not yet in canonical form is either \( \bot \) or contains a constraint \( \forall (A_0; T_1, \ldots, T_n) = \tau \). The ordered condition on patterns prevents the latter (as the second instantiation rule would apply to one of the constraints). Hence, for ordered patterns, rewriting always produces either a canonical unificand or \( \bot \).

Because of the form of the rules, the matching is unitary, and all solutions are equal up to renaming of the \( \alpha_k \)'s. The unique tuple variable that appears in all the \( \tau_j \) can be bound to the tuple \( (\alpha_k) \), and its size bound to the tuple sort.

### 6.4 Related work

The work on dynamics most closely related to ours is that of Leroy and Mauny [17]. Our system can be seen as an extension of their system with “mixed quantification.”

Instead of introducing a typecase statement, Leroy and Mauny merge dynamic elimination with the usual case statement of ML. If we ignore this difference, their dynamic patterns have the form \( QA \) where \( A \) is a type and \( Q \) a list of existentially or universally quantified variables.

For instance,

\[
\forall (A) \exists (F) \exists (B) \exists (G) \left( v:.(A,F,B,G) \right)
\]

is a pattern of their system. The existentially-quantified variables play the role of our pattern variables. The order of quantifiers determines the dependencies among quantified variables. Thus, the pattern above can be rephrased:

\[
\exists (F) \exists (G) \forall (A) \forall (B) \left( v:.(A,F(A),B,G(A,B)) \right)
\]

Writing quantifiers in our patterns explicitly (for ease of comparison), the equivalent pattern in our system is:

\[
\{ F, G \} \left( v:.(P,A,B)T(A,F(P,A),B,G(P,A,B)) \right)
\]

With the same approach, in fact, we can translate all of their patterns into equivalent patterns in our system, preserving the intended semantics.

On the other hand, there does not seem to be a translation from our language to theirs. They have no pattern equivalent to our pattern:

\[
\{ F, G \} \left( v:.(P,A,B)T(A,F(P,A),B,G(P:B)) \right)
\]

because the quantifiers in the prefix of their patterns are in linear order, and hence it is not possible to have the “parallel” dependencies of \( F \) on \( A \) and of \( G \) on \( B \). We can obtain a system intermediate between theirs and ours by leaving tuple variables implicit, and there we would rewrite the pattern above:

\[
\{ F, G \} \left( v:.(P,A,B)T(A,F(A),B,G(B)) \right)
\]

However, we believe that explicit tuple variables are useful, since they allow examples like the applyTwice function:

\[
\text{let applyTwice} = \lambda (dF) \lambda (dxy) \\
\text{typecase } dF \text{ of } \{ F, F' \} (f:F(P)\rightarrow F'(P)) \\
\text{typecase } dxy \text{ of } \{ G, B \} (x,y:F(G(Q)) \times (F(B(Q)))) \\
\text{else } \ldots \\
\text{else } \ldots
\]

This cannot be expressed in our intermediate system, nor in systems with just type quantifiers, such as Leroy and Mauny’s.

### References


Extensions to Standard ML to Support Transactions

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Abstract

A transaction is a control abstraction that lets programmers treat a sequence of operations as an atomic ("all-or-nothing") unit. This paper describes our progress on on-going work to extend SML with transactions. What is novel about our work on transactions is support for multi-threaded concurrent transactions. We use SML's modules facility to reflect explicitly orthogonal concepts heretofore inseparable in other transaction-based programming languages.

1 Revisiting Transactions

1.1 Separation of concerns

Transactions are a well-known and fundamental control abstraction that arose out of the database community. They have three properties that distinguish them from normal sequential processes: (1) A transaction is a sequence of operations that is performed atomically ("all-or-nothing"). If it completes successfully, it commits; otherwise, it aborts; (2) concurrent transactions are serializable (appear to occur one-at-a-time), supporting the principle of isolation; and (3) effects of committed transactions are persistent (survive failures). Systems like Tabs [8] and Camelot [3] demonstrate the viability of layering a general-purpose transactional facility on top of an operating system. Languages such as Argus [4] and Avalon/C++ [2] go one step further by providing linguistic support for transactions in the context of a general-purpose programming language. In principle programmers can now use transactions as a unit of encapsulation to structure an application program without regard for how they are implemented at the operating system level.

In practice, however, transactions have yet to be shown useful in general-purpose applications programming. One problem is that state-of-the-art transactional facilities are so tightly integrated that application builders must buy into a facility in toto, even if they need only one of its services. For example, the Coda file system [7] was originally built on top of Camelot, which supports distributed,
concurrent, nested transactions. Coda needs transactions for storing "metadata" (e.g., inodes) about files and directories. Coda is structured such that updates to metadata are guaranteed to occur by only one thread executing at a single-site within a single top-level transaction. Hence Coda needs only single-site, single-threaded, non-nested transactions, but by using Camelot was forced to pay the performance overhead for Camelot's other features.

The Venari Project at CMU is revisiting support for transactions by adopting a "pick-and-choose" approach rather than a "kit-and-kaboodle" approach. Ideally, we want to provide separable modules to support transactional semantics for different settings, e.g., in the absence or presence of concurrency. Programmers are then free to compose those modules supporting only those features of transactions they need for their application.

1.2 Non-traditional applications

A second problem with existing transactional facilities is that they have been designed primarily with applications like electronic banking, airline reservations, and relational databases in mind. Non-traditional applications such as proof support environments, software development environments, and CAD/CAM systems want transactional features, most notably data persistence, but have different performance characteristics. For example, these applications do not manipulate simple database records but rather complex data structures such as proof trees, abstract syntax trees, symbol tables, car engine designs, or VLSI designs. Also, users interact with these data during long-lived "sessions" rather than short-lived transactions; indeed we can view a "session" itself as a sequence of transactions. For example, during a proof session a user might explore one path in a proof tree transactionally; if the path begins looking like a dead-end the user may choose to abort, backing all the way up to the first node in the path or perhaps to some intermediate node along the way. Also, though multiple users may need to share these data, simultaneous access might be less frequent. For example, proof developers might work on independent parts of a proof tree, perhaps each proving auxiliary lemmas of the main theorem; software developers might modify different modules of a large program. Finally, these non-traditional applications typically support different update patterns. Whereas travel agents make frequent updates to airline reservations databases, we do not expect to make updates as frequently to proofs of theorems saved in proof libraries.

The Venari Project's application domain is software development environments; one specific problem we are addressing is searching large libraries, e.g., specification and program libraries, used in the development of software systems. We imagine the scenario in which a user searches a large library for a program module that "satisfies" a particular specification. We might wish to perform each query as a transaction, for example, to guarantee isolation from any concurrent update transaction or to abort the query after the first n modules are returned.

Our effort to support a "pick-and-choose" approach for transactions has the advantage of providing us with a way to take performance measurements on different combinations of our separable modules. We have the potential to do different kinds of performance tuning for the non-traditional applications we hope to support.

1.3 Contributions of this paper

SML's modules system lets us cleanly exemplify our "pick-and-choose" approach. In the case of single-site, single-threaded, nested transactions, we support separately the persistence and undoability prop-
erties of transactions in terms of two different modules; we then compose them to build a module for transactions [6]. We reported on this work at the Pittsburgh 1991 ML workshop. Also, along with others at Carnegie Mellon, we have separately designed and built a threads package for SML/NJ [1]. We reported on this work at the Edinburgh 1990 ML workshop. 1

This paper reports on our progress for combining our support for threads and that for transactions. We address concurrency, in two ways: making an individual transaction multi-threaded and allowing multiple transactions to run concurrently. To the transaction and database community, our work is novel because it is the first to cast within a programming language a model of computation that supports multi-threaded transactions. To the programming language community, our work is among the few to extend the functional programming paradigm to support a traditionally imperative feature. To the SML community, our application of the modules facility should be of particular interest.

In the rest of this paper, we motivate the desire to keep threads and transactions as separate control abstractions (Section 2), give a snippet of the SML programmer’s interface to multi-threaded concurrent transactions (Section 3), describe some details of our design (Section 4), discuss what features of SML helped in our design and implementation effort (Section 5), and close with a list of future work (Section 6).

We emphasize that this paper describes on-going work. We are taking a pragmatic, bottom-up approach; by prototyping individual features (e.g., persistence, undoability, read/write locking, nesting, threads, and transactions) incrementally and then combining them in various ways, we can explore a rich design space. This paper focuses on the hardest combination: threads and transactions. (Examples of other reasonable combinations are “multi-threaded persistence” and “multi-threaded undoability.”) Thus, for now we are concerned with providing efficient enough run-time mechanism to give systems builders flexibility in deciding policy. Although the design described in Section 4 does reflect one particular policy, our runtime mechanism is general enough to support alternate policies. Finally, we have not thought greatly about the “ideal” programming interface to provide the SML end-user, but look forward to designing one in the near future.

2 Keeping Threads and Transactions Separate

In languages like Argus and Avalon, a single thread of control is associated with each transaction. But threads and transactions are orthogonal control abstractions. So, we would like to relax the restriction of identifying threads and transactions by allowing multiple threads of control to execute within, and on behalf of, a single transaction.

Figures 1a and 1b depict the traditional model, where we use a wavy line to denote a thread and a box to denote a transaction; time moves from left to right. Figure 1a shows a single thread executing, first entering a transaction and then leaving successfully (i.e., committing). Figure 1b shows two single-threaded transactions executing concurrently. Figure 1c depicts our new model where multiple threads execute within a single transaction. And finally, Figure 1d depicts multi-threaded concurrent transactions, the “composition” of Figures 1b and 1c. The goal of Venari’s version of SML is to support Figure 1d through module composition.

1For this paper to be somewhat self-contained, we include the cited Pers, Undo, Trans, and Threads signatures in Appendices A and B.
Figure 1: Threads and transactions are separate control abstractions.
2.1 Why have multiple threads within a transaction?

The most compelling argument for supporting multiple threads within a transaction is modularity. Consider the following kinds of multi-threaded programs: (1) a search procedure that uses multiple threads to find program modules satisfying a specification, returning when the first one is found; (2) a procedure with benign side effects, e.g., rebalancing a B-tree or doing garbage collection, that executes in the background of the main computation; (3) a netnews server that uses multiple threads to minimize latency.

We would like to be able to run such a multi-threaded program from within a transaction without having to modify the source code. We would like to treat the program as a black box, reuse it in its entirety, but have its effects done transactionally (i.e., atomic, serializable, and persistent). Without being able to simply “wrap” a transaction around the program, we are forced to recode each separate thread as a concurrent subtransaction of a top-level transaction. This violates one aspect of modularity since the entire program has to be recoded.

2.2 Why have multiple threads at all?

Concurrent transactions have to be serializable. Thus, by definition, we can view transactions as happening one after another. On the other hand, threads are often used for two-way communication through shared, mutable resources (e.g., refs). If we identify each thread with a single transaction, then we can no longer do two-way communication between threads. For instance, assuming we associate each thread with a transaction, then Figure 2a shows thread/transaction A and thread/transaction B executing concurrently. Transaction semantics require that the effects of A and B executing concurrently are the same as that of either A executing first followed by B (Figure 2b), or vice versa. Suppose A sends a message to B and B wants to acknowledge A; we cannot put A’s execution before B (since A will never get the acknowledgment) nor can we put B before A (since B will never get the message). Thus if we want to support two-way communication between processes, we need to support multiple threads independent of transactions.

Another argument for supporting both threads and transactions as orthogonal concepts is performance. In existing transactional systems, the runtime cost of creating and managing a transaction (“heavyweight” process) is not the same as that for a thread (“lightweight” process). Transactions require runtime mechanism to support protocols for locking, logging, committing/aborting, and crash recovery. There are cases when parallelism is desired without the performance overhead of transactions. Again, even if we were to recode one of our example multi-threaded programs with transactions, we probably do not want to incur the cost of making each thread a transaction.

In short, transactions provide features that threads do not: persistence, undoability, isolation of effects, atomicity of a sequence of operations, and crash recovery. Threads provide functionality, e.g., two-way communication, and performance benefits that transactions do not.
Figure 2: Transactions are serializable.
3 Design Overview

As in our design for single-threaded transactions for SML [6], if f is a function applied to some argument a, then to execute:

\[ f(a) \]

in a transaction, we want programmers to be able to write:

\[(\text{transact } f)(a)\]

or more probably,

\[(\text{transact } f)(a) \text{ handle } \text{Abort} \Rightarrow \text{[some work]}\]

Here f might be multi-threaded. Informally, the meaning of calling f with \text{transact} is the same as that of just calling f with the following additional side effects: If f returns normally, then the transaction commits, and if it is a top-level transaction, its effects are saved to persistent memory (i.e., written to disk). If f terminates by raising the exception Abort, then the transaction aborts and all of f’s effects are undone. Through SML’s exceptional handling, in the case of an aborted transaction, the programmer has control of what to do such as clean-up and/or reraising Abort. Note that we support the usual model for subtransactions: the persistence of a child’s effects is relative to the commit of its parent and aborting a child does not imply the abort of its parent.

We have implemented the interfaces shown in Figure 3. We use standard two-phase read/write locks to ensure serializability among concurrent transactions. We use Moss’s locking rules for nested concurrent transactions [5].

Two items of note are visible through these interfaces. First, the \text{TRANSACTION} signature shows the clear separation between threads (the substructure Thread.System) and transactions (the substructure Trans). The functor header additionally shows how we have achieved modularization of our support: concurrency within a transaction is packaged in TRANS.THREAD; concurrency among transactions, in RW.LOCK; transaction undoability, in UNDO.

Second, we guarantee the principle of isolation for transactions by making use of “safe” refs [9] (and correspondingly “safe” arrays). In the context of just threads, a normal SML ref is unsafe, while a ref protected by a mutex is a safe ref. In the context of transactions, a ref protected by only a mutex is an unsafe ref, while a ref protected by both a mutex and a read/write lock is a safe ref. A read or write of a safe ref will fail unless the thread (transaction) holds the mutex (read/write lock) of the ref. Thus, it is impossible to violate the isolation principle if the programmer uses only safe refs.

\(^2\)Appendix B shows parts of the THREAD.SYSTEM and other relevant signatures; see [1] for a discussion of threads in SML.
signature TRANSACT =
  sig
   structure Trans :
     sig
      exception Abort
      val transact : ('a -> 'b) -> 'a -> 'b
      val abort_top_level: unit -> 'a
      val abort: unit -> 'a

      eqtype rw_lock
      val rw_lock : unit -> rw_lock
      val acquire_read : rw_lock -> unit
      val acquire_write : rw_lock -> unit
   end

   structure SRef : SREF
   structure SArray : SARRAY

   structure Thread_System : THREAD_SYSTEM

   sharing type SRef.lock = SArray.lock = Trans.rw_lock
   sharing type SRef.uref = Thread_System.SRef.sref
   sharing type SArray.uarray = Thread_System.SArray.sarray
  end

functor Transact (structure TT TRANS-THREAD
   structure RW : RW_LOCK
   structure SRef : SREF
   structure SArray : SARRAY
   structure Undo : UNDO
   sharing type SRef.lock = SArray.lock
   sharing type SArray.lock = RW.T
   sharing type SRef.uref = TT.TS.SRef.sref
   sharing type SArray.uarray = TT.TS.SArray.sarray)
  : TRANSACT = struct ... end
4 Design Details

4.1 Simplifying assumptions

To simplify our model, and hence our design, we assume that there is exactly one thread that enters a transaction and exactly one that leaves a transaction. We do not lose any generality since we can always immediately fork off multiple children upon entry and we can always force all threads to join into one upon exit. Second, again without loss of generality, we will assume that conceptually the thread exiting is the same as that entering; we call this the transaction's root thread. Finally, we assume no mutexes are held before a transaction begins. We make this assumption so we do not have to reacquire locks that were held upon entering a transaction in case an abort occurs. Doing so could cause a deadlock: Suppose the entering thread t holds a lock and then releases it sometime during the transaction. A thread s outside the scope of the transaction then acquires it. If the transaction now aborts, and we are to undo all of its effects, including reacquiring the lock, we may deadlock if s is waiting to acquire some other held lock.

4.2 Per transaction state

Just as there is per thread state [1], we assume there is per transaction state. This state includes four pieces of separable information:

- The data objects accessed by the transaction. Since the order of modifications to these data objects is important with respect to abort, we call this information the (data) undo list.

- The set of mutex locks held by threads within a transaction. We call this information the mutex lock set.

- The set of read and write locks held for the duration of the transaction. We call this information the read-write lock set.

- The set of threads running on behalf of the transaction.

The first piece of information (data state) is separable from the other three (synchronization state) which we need to maintain because of concurrency due to threads.

4.3 Who commits and who aborts?

Our design gives the root thread the privilege of committing the transaction and the responsibility of knowing when it is safe to do so. However, for abort, any thread may encounter a state in which the thread cannot back out of and knowingly wish to cause the abort of the entire transaction; the root thread need not be the only thread to determine that an abort is necessary. Thus, rather than requiring such a thread to communicate with the root thread who could then cause the abort, we permit any thread to cause an abort.

3 We could relax this assumption by letting threads pass a “baton” among each other, where the baton’s flow of control would reflect that of the root thread, but this relaxation is unnecessarily general.
4.4 What happens upon commit?

The effect of a commit is to preserve all data state changes made by the committing transaction. Upon commit, we do the following:

1. Stop all other active threads running on behalf of the transaction so they do not continue to modify state;
2. If the transaction is top-level, throw away the data undo list since we do not need to save the old data values; otherwise, anti-inherit the list to its parent.
3. Release all mutex locks held by non-root threads running on behalf of the transaction.
4. Anti-inherit all read/write locks to its parent [5].
5. If the transaction is top-level, save the state of persistent memory.
6. Exit the transaction and continue processing.

4.5 What happens upon abort?

A transaction may voluntarily abort or be involuntarily aborted (e.g., due to a system crash). Following our semantics for single-threaded transact [6], we mask any exception as an abort. Moreover, we treat any unhandled exception as an abort. The effect of an abort is to undo all changes to the data state made by all threads executing on behalf of the transaction. Upon abort, we do the following:

1. Stop all other active threads running on behalf of the transaction so they do not continue to modify state;
2. Follow the undo list backwards, rewriting all old data values.
3. Release all mutex locks held by threads running on behalf of the transaction.
4. Release all read/write locks.

It is critical that we first undo the data values, then release mutexes, and then release read/write locks. Data are protected by mutexes; read/write locks are implemented using them. If we were to release mutexes before undoing all data values, then a thread may modify a data object after the old value gets rewritten. In order to release a read/write lock, we need to be able to acquire other mutexes; if we were not to release mutexes before read/write locks, we could end up in a deadlock situation.

5 Where SML Made a Difference

The SML modules facility is key to our “pick-and-choose” approach. It lets us explicitly reflect the inherent orthogonality of concepts like persistence, undoability, multi-threading, and transactions by letting us define separate structures for each and then functors that compose them. The Transact functor that builds a structure for multi-threaded transactions is one example (Section 3). When we prototyped our implementation for single-threaded, non-concurrent transactions (Appendix A), we also
used a functor parameterized over Pers and Undo structures, which respectively provide persistence and undoability.

We also parameterized the Thread.System structure itself so that the programmer can pick-and-choose among separable support for persistence, undoability, and multi-threading. The Fox Project at CMU, for example, needs only multi-threading; it does not have to use a separate threads module, but rather it just has to apply the Thread.System functor to Undo and Pers structures that are essentially "no-ops." Moreover, it is to SML's credit that modifying the original Thread.System structure to work with Undo required only two lines of additional code: to "turn off undo" while doing a thread operation (e.g., acquiring a mutex) and to "turn it back on" when the operation is completed.

Another way we are able to exploit the modules facility is in code reuse. For example, we use only one functor to implement both kinds of safe refs, that for just threads and that for transactions (q.v., sharing type SRef.u.ref = TT.TS.SRef.sref of Figure 3).

Having first-class functions in SML lets us easily support first-class transactions. This view of transactions is a radical departure from the more traditional view taken by other transaction-based programming languages. Programmers in Camelot, Argus, and Avalon write constructs like begin transaction ... end.transaction to bracket transaction boundaries and cannot treat the compound statement as a value.

We relied on the "mostly functional" nature of SML in our implementation of undoability and persistence. For example, our implementation for undoability keeps a log of all modifications to the store and the old values originally assigned to the modified locations. For traditional imperative languages where modifications to the store would be frequent, maintaining and replaying such logs would be expensive. Such a log is inexpensive to maintain for SML since we can assume mutations are rare.

Though we greatly benefit from SML's static typing, one place where we need dynamic types is in our support for persistence. Our interface allows us to add bindings between identifiers and values to a persistent environment; SML cannot statically determine whether the type of the value returned by a subsequent retrieve (e.g., upon startup of a new SML session or upon crash recovery) on some identifier is the same as the type of the value when it was initially bound.

In summary, SML is a nice vehicle with which to express separable concepts. Though SML may not be the natural language of choice in which to support transactions, it is a natural language of choice for the non-traditional applications of transactions that we have in mind. Many of CMU's projects that involve reasoning about programs, Edinburgh's LEGO "proofs-as-programs" system, Cornell's NuPrl system, and AT&T Bell Labs's proof support environment all use SML as their implementation language. These applications need some, if not all, transactional features like data persistence, concurrency control, checkpointing, backtracking, and crash recovery. We hope to provide these potential users with a set of SML modules that they can use in a "pick-and-choose" fashion.

6 Status and Future Work

We have a working prototype of all the interfaces given in this paper, but much work remains:

- **Language design:** As mentioned in the introduction we have yet to do a design of an SML end-user's interface for multi-threaded concurrent transactions. We are also still exploring different policies that our mechanisms can support. We may export different end-user interfaces, each reflecting a different policy.
• **Semantics**: We have been negligent in working out any formal semantics for our extensions to SML. Some of the challenges specific to SML include a semantics for undoability and a semantics for the interactions between callcc and transact; specific to transactions, a model of computation and meaning of correctness for multi-threaded concurrent transactions.

• **Implementation**: After our prototype becomes stabler, we intend to build sample applications and perform experiments to measure the costs of our extensions.

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**References**


Appendix A: Persistence and Undoability

Figure 4 illustrates persistence and undoability as orthogonal concepts; we compose them to form single-threaded, non-concurrent, nested transactions. In Figure 4a, as a thread executes, a call to persist has the effect of saving all values reachable from a persistent handle to disk. In Figure 4b, a call to checkpoint has the effect of "remembering" the store at the point of call; a call to restore has the effect of undoing the effects on the store, reverting back to that at the (dynamically) last call to checkpoint. Finally, Figure 4c shows how we compose checkpoint with persist to give us transaction commit; checkpoint with restore to give us transaction abort. The signatures for persistence, undoability, and transactions that follow are unfortunately slightly different from that described in [6], but do reflect the current working version of our implementation. The primary difference between the TRANSACT signature here and its analogue in Figure 3 is the absence of an interface for read/write locks, which is not needed in the absence of concurrency.

signature PERS = sig
  exception INIT_FAILED
  exception PERSIST_FAILED
  val init: string * string * bool -> unit
  val persist: ('a->'b) -> 'a -> 'b

  type identifier
  exception UnboundId
  val bind: identifier * 'a -> unit
  val unbind: identifier -> unit
  val retrieve: identifier -> 'a
end

signature UNDO = sig
  exception Restore of exn
  val checkpoint: ('a -> 'b) -> 'a -> 'b
  val restore: exn -> 'a

  val exn2restore: ('a -> 'b) -> 'a -> 'b
  val restore2exn: ('a->'b) -> 'a -> 'b
  val restore_on_exn: ('a -> 'b) -> 'a -> 'b
end

signature TRANSACT = sig
  val transact: ('a -> 'b) -> 'a -> 'b

  exception Abort
  val abort_top_level: unit -> 'a
  val abort: unit -> 'a
end
Figure 4: Persistence + Undoability = Transactions
Appendix B: Threads

Our Threads interface and parts of the TRANS THREAD and THREAD SYSTEM signatures:

signature THREAD = sig
  val fork : (unit -> unit) -> unit
  val exit : unit -> unit

  type mutex
  val mutex : unit -> mutex
  val with_mutex : mutex -> (unit -> 'a) -> 'a

  type condition
  val condition : mutex -> condition
  val with_condition : condition -> (unit -> 'a) -> 'a
  val signal : condition -> unit
  val broadcast : condition -> unit
  val await : condition -> (unit -> bool) -> unit
  val vwait : condition -> (unit -> 'a option) -> 'a

  exception Undefined
  type 'a var
  val var : unit -> 'a var
  val get : 'a var -> 'a
  val set : 'a var -> 'a -> unit

end

signature TRANS THREAD = sig
  structure TS : THREAD SYSTEM
  structure TransID : sig ... end
end

signature THREAD SYSTEM = sig
  structure Thread : THREAD

  structure SRef : sig ... end
  structure SArray : sig ... end

end
Programming Images in ML

Introduction

In this paper, we describe a library that provides basic types and functions to produce graphical documents in ML. This library is implemented on top of CAML [9] and CAML Light [8] in a functional style [7].

The graphical model is basically that of PostScript [1]. Various objects can be defined on the infinite cartesian plane and arbitrarily scaled, translated and rotated by application of linear transformations. Moreover, high level primitives allow for various combinations of objects.

The main difference with PostScript is that graphic objects are represented by a data structure and has a ML type. A type “picture” is used to represent all printable objects. Pictures have a “frame” and possibly a set of named “handles” that are used for combination operations. Pictures are defined from more basic objects such as geometric elements (polygonal lines, circle arcs and Beziers curves), texts and bitmaps. All operations defined on these types are purely functional except for pixel editing in bitmaps.

Printing is obtained via a translation to PostScript. The philosophy of this translation has been to delegate as much work as possible to PostScript. In particular, the application of linear transformations to pictures is delegated to the PostScript interpreter. This has two advantages: the efficiency of PostScript interpreters is fully used and the sharing involved in the ML representation of pictures is preserved as much as possible.

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The main application we foresee for this library is the production of documents involving integrated use of ML and PostScript or ML, \textsc{\LaTeX} and PostScript as in the text you are presently reading. In particular, technical texts describing applications written in ML (e.g., proof systems or abstract machines) can be easily decorated with visual representations of ML objects involved in computations (cf. figure 12 showing pictures from Y. Lafont [6] and P. Crégut [4]). Visualization of ML objects can also be used interactively by running an ML window and a PostScript window concurrently. Adding graphical attributes to ML types could lead to interesting debugging tools.

As we all know, functional languages have an expressive power which facilitates the programming activity. It is therefore not surprising that our library provides a way of producing images which is much more pleasant than using the PostScript language itself.

The system we have obtained so far is only a basic layer for producing images in ML. We think it is a good basis for writing technical pictures, described by programs as some of our examples show. We plan to develop further layers to do so.

The paper gives an overview of the system, followed by application examples which reflect some of its possibilities. The last section describes some possible further developments.

1 Overview of the system

1.1 Basic concepts

Each graphic notion has a corresponding ML type. Objects that can be made visible on a graphic device are of type \textit{picture}. A picture always has an absolute position on the infinite cartesian plane. The user can use this fact if it is convenient for her/him or completely forget about it by using more abstract constructions. Pictures can be arbitrarily translated, rotated and scaled. Some pictures can also be created using non-linear transformations (cf. figure 11).

Pictures are produced by grouping more elementary pictures in various ways. Basic constituents of pictures are sketches, texts and bitmaps. A sketch is a sequence of geometric elements such as be lines, arcs and curves. A sketch can be transformed into a picture by choosing a linestyle and a color to draw its constituents or a fillstyle and a color to draw its interior. It can also be used to clip part of a picture. Before being used to produce pictures, sketches can also be arbitrarily grouped and transformed.

1.2 Sketches

A sketch is basically a sequence of geometric elements. It corresponds to what is called a “path” in the PostScript terminology.

The basic constituents of sketches are geometric elements. A geometric element is either a polygonal line represented by a sequence of points or a circle arc represented by a center, a radius and two angles or a Beziers curve represented by a start point, two control points, and an end point. The corresponding CAML types are the following:

\begin{verbatim}
  type point = {xc:float;yc:float};;
  type geom_element =
\end{verbatim}
Given the points $A=(1.1), B=(1.3), C=(3.3), D=(3.1), E=(1.8), F=(2.4), G=(4.9), H=(5.8)$ and $I=(7.4)$, the expressions $\text{Seg} [A;B;C;D;A]$, $\text{Curve}(E,F,G,H)$ and $\text{Arc}(I,2,30,290)$ correspond to the three elements that are drawn with thick lines in figure 1.

Figure 1: Geometric elements

Sketches are represented by a CAML type sketch which is used as an abstract type. The representation of type sketch involves lists of geometric elements together with additional information such as frame and interface information. The basic building function for sketches is:

make_sketch : geom_element list -> sketch

1.3 Painting Information

Sketches are made of pure lines. To be visualized as images, they require painting information which are represented by the types linestyle, fillstyle and color.

type linestYLE = {linewidthLS:float; linecapLS:linecap; linejoinLS:linejoin; dashpatternLS:float list};
type fillstyle = Nzfill | Eofill;;
type color = Rgb of float * float * float
           | Hsb of float * float * float
           | Gra of float;;

We do not detail here the meaning of these types which refer exactly to PostScript notions.
1.4 Pictures

The type picture is used for all visual objects in the system. Pictures can be built from sketches using painting information. As shown later, they can also be built from bitmaps and texts.

Here are the two functions that make pictures from sketches:

\[
\text{make\_draw\_picture} : \text{linestyle} \times \text{color} \rightarrow \text{sketch} \rightarrow \text{picture} \\
\text{make\_fill\_picture} : \text{fillstyle} \times \text{color} \rightarrow \text{sketch} \rightarrow \text{picture}
\]

Pictures can be “put together” using functions:

\[
\text{join\_pictures} : \text{picture} \rightarrow \text{picture} \rightarrow \text{picture} \\
\text{join\_picture\_list} : \text{picture list} \rightarrow \text{picture}
\]

The arguments of these functions are taken from left to right. Each new picture can cover previous ones partially or totally. Figure 2 shows examples of what can be done using a sketch representing letter P. The left image is a draw picture, the middle one is a fill picture and the right one is a superposition of the two.

![Figure 2: Three pictures obtained from the same sketch](image)

1.5 Picture transformations

Transformations are normally functions of type \text{point} \rightarrow \text{point}. However, most useful transformations are linear transformations represented by \(3 \times 3\) matrix which have shape:

\[
\begin{pmatrix}
    m_{11} & m_{12} & m_{13} \\
    m_{21} & m_{22} & m_{23} \\
    0 & 0 & 1
\end{pmatrix}
\text{operates on vector}\begin{pmatrix}x \\
y \\
1\end{pmatrix}
\text{representing point \((x, y)\).}
\]

Users normally build linear transformations using library functions such as:

\[
\begin{align*}
\text{translation} : \text{float} \times \text{float} \rightarrow \text{transformation} \\
\text{rotation} : \text{point} \rightarrow \text{float} \rightarrow \text{transformation} \\
\text{scaling} : \text{float} \times \text{float} \rightarrow \text{transformation} \\
\text{compose\_transformation} : \text{transformation} \rightarrow \text{transformation} \rightarrow \text{transformation} \\
\text{inverse\_transformation} : \text{transformation} \rightarrow \text{transformation}
\end{align*}
\]

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Transformations are applied to pictures using function

\[
\text{transform\_picture} : \text{transformation} \rightarrow \text{picture} \rightarrow \text{picture}
\]

Figure 3 describes the effect of applying transformations T1= translation (2,-9), T2= scaling (0.5,0.5) and T3= rotation xc=-2;yc=-2 60 to a basic picture.

![Figure 3: Transformations T1,T2 and T3](image)

1.6 Frames

The CAML graphic system maintains a frame information for pictures. A frame is a rectangle with sides parallel to the axes which contains the picture. It can be made to be the smallest rectangle containing the object but the user can also choose to determine the frame of an object by himself and under his own responsibility for instance to add blank space around an object. Frames provide the basis to define various operations such as rotation of an object around its center (defined as its frame center), horizontal and vertical flip operations, changing the size of an object to fit it in a different frame or putting an object besides or over another object adjusting their frame width and frame height.

Typical functions using frames explicitly are:

\[
\text{frame\_center} : \text{frame} \rightarrow \text{point}
\]
\[
\text{frame\_to\_frame\_transform} : \text{frame} \rightarrow \text{frame} \rightarrow \text{transformation}
\]
\[
\text{picture\_frame} : \text{picture} \rightarrow \text{frame}
\]
\[
\text{extend\_picture\_frame} : \text{extension} \rightarrow \text{float} \rightarrow \text{picture} \rightarrow \text{picture}
\]

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The functions `extend_picture_frame` and `force_picture_in_frame` are used under user's responsibility to give arbitrary frame to object. The function `fit_picture_in_frame` involves scaling and translating a picture to make it fit exactly in a given frame. The transformation that is used to do so is accessible to the user via function `frame_to_frame_transform`. The figure 4 shows the result of fitting a given picture in a given frame.

![Our Example](image1) ![A frame](image2) ![Example fit into frame](image3)

**Figure 4: Fitting a picture into a given frame**

Some local picture transformations are computed using the frame such as the following ones:

- `rotate_picture : float -> picture -> picture`
- `vflip_picture : picture -> picture`
- `hflip_picture : picture -> picture`

Figure 5 shows the effect of these functions on our favorite example:

![rotation by 45](image4) ![vertical flip](image5) ![horizontal flip](image6)

**Figure 5: Transformations using frames**

1.7 Texts

Text pictures are strings of characters that will be displayed in a given font style and a given size. They are built with their bottom left corner at the origin of the coordinate system but then they can of course be moved to any position.
The basic functions for producing text pictures are:

\[
\begin{align*}
\text{make_font} & : \text{font_style} \rightarrow \text{float} \rightarrow \text{font} \\
\text{make_text_picture} & : \text{string} \rightarrow \text{font} \rightarrow \text{color} \rightarrow \text{picture}
\end{align*}
\]

An extensive list of font styles exists and the user can have access to the width of any string in any font. Using this information, one can for instance write text on a circle in the following way:

```ocaml
#let Pi = acos (-1.0);;
Pi : float
Pi = 3.14159265358979

#let circletext fnt str center radius =
#  let l = text_width fnt str in
#  if l >= 2.0*Pi*radius
#    then failwith "text is longer than circumference"
#    else let start_angle = (Pi/2.0) + 1/(2.0*radius)
#      and char_list = explode str in
#      let angle_list = start_angle::(rev(tl (snd
#        (it_list (fun (a,l) w -> let aa = a-w/radius
#          in (aa, aa::l))
#        (start_angle,[]))
#        (map (text_width fnt) char_list))))
#      and place_char (s,a) =
#        let T1 = rotation center ((a-Pi/2.0)*180.0/Pi)
#        and T2 = translation (center.xc,center.yc+radius)
#        in transform_picture (T1 CT T2)
#        (make_text_picture s fnt (Gra 0.0))
#      in join_picture_list
#      (map place_char (combine (char_list,angle_list)));;

The result of applying circletext with center (5.5) and radius 3 is shown in figure 6.

![Figure 6: A circular text]

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1.8 Bitmaps

A bitmap is basically a two dimensional array of pixels. It has a height, a width and a depth. The depth is the number of bits used for each pixel. Possible values are 1, 2, 4 or 8.

Bitmaps can be created from scratch using functions

```
create_bitmap : int -> int -> int -> bitmap
set_pixel : bitmap -> int -> int -> int -> char
```

However, they are usually read from a string representation obtained through a digitalizer by function:

```
read_bitmap : int -> string -> bitmap
```

whose first argument is the depth.

Functions are provided to uniformly modify a bitmap or extract a sub-bitmap such as

```
map_bitmap : (int -> int) -> bitmap -> bitmap
sub_bitmap : bitmap -> int * int -> int * int -> bitmap
```

Figure 7 shows an example of a bitmap of depth 1 and its inverted image.

```
camel.bmp  map_bitmap (fun 0-> 1; 1->0) camel.bmp
```

Figure 7: A bitmap transformation

1.9 Picture Composition using frames

Frames can be used to adjust pictures such that their frames fit nicely together either side by side or one on top of the other. Functions BPICT and OPICT make a new pictures by putting two pictures side by side or one on top of the other. If necessary, the second argument is scaled so that its frame gets same height or width as that of the first argument. Other functions perform similar actions but do not scale the second argument. The two arguments are then aligned in specified ways.

Using these composition functions, the users can forget completely about the cartesian plane in which pictures are built. Only the relative sizes of pictures are relevant.

This technique of picture composition is nicely exemplified by the construction of an Escher picture called “Square limit”. A programmed version of this picture has
been given by Henderson in [5]. A detailed account of Henderson's approach is also given in Course Notes by Cousot [3]. The version presented here uses the same basic pictures but builds the final picture in a different way.

The building blocks of the final picture are four basic pictures $A,B,C,D$ and three functions named trio, quartet and cycle.

Here are the four basic pictures:

![Picture A](image1)
![Picture B](image2)
![Picture C](image3)
![Picture D](image4)

The three basic functions are defined in the following way:

```ml
#let rot = rotate_picture 90.0;;
rot : picture -> picture
#let trio(p1,p2,p3) = (p1 BPICT p2) OPICT p3;;
trio : picture * picture * picture -> picture
#let quartet (p1,p2,p3,p4) = (p1 BPICT p2) OPICT (p3 BPICT p4);;
quartet : picture * picture * picture * picture -> picture
#let cycle p = (p BPICT (rot (rot (rot p))))
  OPICT ((rot p) BPICT (rot(rot p))));;
cycle : picture -> picture
```

Here is a description of what these functions do:

![Trio](image5)
![Quartet](image6)
![Cycle](image7)

The CAML definition of the picture construction is:

```ml
let small = scale_picture (0.5,0.5);;
let square_limit n (P,Q,R,S) =
  let TT=quartet(P,Q,R,S)
  and U=cycle (rot Q)
```

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in
let step(C,L,T) =
     (quartet(small C,
              small(L BPICT T),
              small((rot T)OPICT(rot L)),
              ,UU),
     trio(small L,small T,rot TT),
     trio(small L,small T, TT))
and final_step(C,L,T) =
    quartet(small C,small L, small(rot T),rot Q)
in
    cycle(final_step(iterate step n (TT,rot TT,UU)));;

Here are TT and UU for basic picture A,B,C,D:

The picture corresponding to the original Escher etching is
square_limit 2 (A,B,C,D). It is shown together with its structural description in
figure 1.9.

1.10 Another form of picture composition

Pictures can be given handles in order to combine them in specific ways to obtain new
pictures. A handle is an oriented segment defined by two points. The simplest case is
when pictures have one input handle and one output handle. Pictures can also have 
sets of named handles in input and output.

The function `APICT` combines two pictures by transforming the second in such a way 
that the output handle of the first one coincides with the input handle of the second 
one. The input and output handles of the result are the input handle of the first one 
and the output handle of the second one.

Figure 8 demonstrates the use of handles. We start with a square bitmap with side 
equal to one. It has two handles. The input handle is the segment `((0,0),(1,0))` at the 
bottom of the picture. The output handle is the segment `((1,1),(1,1/::-φ))` where φ is 
the golden number `(1 + √5)/2`. The result is obtained by applying to it the function 
gold_spiral defined by:

\[
\text{let rec gold_spiral P = function 0 -> P
| n -> (gold_spiral(n-1)) APICT P;;}
\]

![Figure 8: Golden Camels](image)

1.11 Displaying pictures

The display functions produce a PostScript translation for a picture which can be 
directed to a file. These functions can perform a translation and a uniform scaling 
of the picture in order to make it centered on the page with reasonable size. So the 
user does not have to worry about the actual position and size of its image in order to 
visualize it.

2 Application examples

2.1 Drawing binary trees

Drawing pictures of binary trees that are pleasant to look at requires some computa-
tion. The main constraint to satisfy is that subtrees should be not overlap. Another 
constraint, almost equally important, is that space should be rather uniformly occupied 
i.e. given two subtrees with the same father, the respective space to allocate to each 
of them depends on their size and shape. Figure 9 shows two rather different kinds of 
binary trees.

The design principles we have adopted for drawing binary trees are the followings:
At each level of a binary tree, the distance between brother nodes should be constant.

The distance between brother nodes at level \((n+1)\) should at most equal to the distance at level \(n\).

Two brother subtrees should be drawn in such a way that at each level, the distance between the rightmost node of the left subtree and the leftmost node of the right subtree should be at least equal to the standard distance between two brother nodes at that level.

These constraints are taken into account by a function `compute_coef_list` which computes for each binary tree a list of coefficients which indicate the ratio that should be adopted between the distance between two brother nodes at level \((n+1)\) and the distance between two brother nodes at level \(n\). Given this list of coefficients, the drawing of a binary tree becomes straightforward. The drawing function is a standard recursive function on binary tree which has as a parameter a function to draw nodes. Different functions can therefore be used to draw nodes. For instance, if trees are AVL trees, it is possible to indicate in each node whether the subtree corresponding to each node is balanced, or heavier on the left or heavier on the right as shown in figure 10.

![Figure 9: Two examples of binary trees](image_url)

![Figure 10: An AVL tree](image_url)
2.2 Escher’s Circle Limit III picture

The picture shown in figure 11 was programmed by students at École Normale Supérieure [2]. It uses knowledge from hyperbolic geometry and group theory.

The circle that contains the picture is the Poincaré representation of the hyperbolic plane. The Escher design is based on a paving of the hyperbolic planes using an isometry group applied to a unique basic picture which is a fish. This group has three generators. By adding explicitly their inverses, it is possible to obtain a canonical rewriting system for checking equality of its elements. It is therefore possible to generate non redundant sets of elements of the group. The picture is obtained using a finite non redundant subset.

The isometries involved correspond in the Poincaré model to homographies of the form:

\[ h_{a,\lambda} : z \mapsto \frac{\lambda z + a}{1 + \bar{a}z}, \]

where \( a \) and \( \lambda \) are complex numbers verifying \(| a | < 1 \) and \(| \lambda | = 1 \).

These non linear transformations have to be computed by ML since PostScript does not know how to handle them. The finite set of fishes that is retained for the final picture are those whose size is greater than a given bound.

The key function is extremely simple. It takes as arguments a minimal bound \( d \) for the size and a list of transformations:

```ml
let rec escher d =
  let ok x = module (app (calc x) q1) < d in
  fun [] -> black_circle
  | (x::l) -> if ok x then (escher d) PICT (poisson x)
                else (escher d l);;
```

Figure 11: Circle Limit III
2.3 Others contributions

We show two works realized at LIENS by P. Créguet and Y. Lafont (cf. figure 12).

P. Créguet gave a trace in his paper [4], for his abstract machine, to reduce $\lambda$-terms. The figure 12 shows the reduction from $\lambda t. (\lambda u. u (\lambda v. u)) (\lambda z. x) t$ to $\lambda t (\lambda v. t)$. The conventions used in the drawing are:

- applications are represented by a white circle,
- bound variables by a black circle with the De Bruijn's index,
- free variables by a white circle with the nesting level,
- closures by a white box with the term, its environment, the nesting level and the nesting level of the closure,
- the current state by a black box like a closure with a stack as third term.

![Figure 12: Trace execution and Rewriting rules](image)

For some experiments in 2-dimensional symbolic computation, Y. Lafont is developing a software in the functional programming language CAML using this graphic library. It was used to check the confluence of his twelve rules (only few seconds for checking, a bit more for printing). The fifty-six critical pairs are shown in his paper annex [6] under graphical form.
3 Possible Developments

The library presented here offers the possibility of representing complex images as ML values of type picture and gives functions to operate on this type. Basically, it incorporates in ML the possibilities of PostScript with an added value offered by the functional style.

Typical applications are functions of type user_type -> picture which give a visual representation for conceptual objects such as for instance binary trees. It is clear that many objects defined by ML users have a visual counterpart that is much more readable and informative than what the standard value printer produces. Associating visualization functions to ML types could be an important debugging aid. It could also greatly facilitate the production of research papers describing systems implemented in ML. The present system is too raw and requires too much effort from the user to produce easily images from ML types. We shall therefore investigate the possibility of building appropriate tools to help him/her in this task. We shall also investigate the possibility of producing informations about the ML system in a graphic way.

Another possibility of application is the production of complex technical pictures. The present situation in this domain is rather frustrating. The production of technical texts has been greatly facilitated by the existence of \TeX but no similar facility exists for pictures. We think that a functional description of pictures is a good basis for improving this situation.

References


Distributed Programming with
Asynchronous Ordered Channels in Distributed ML

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ABSTRACT

Distributed ML (DML) is an extension of Standard ML for reliable distributed programming. This paper motivates the choice of port groups and asynchronous multicast as DML’s distributed communication primitives [Krumvieda 1991].

Multicast is an important tool for parallelism and fault tolerance in distributed programming. It is useful whenever groups of processes must cooperate on some task, or receive the same piece of information. Writing distributed programs comprising many processes is made difficult by the abundance of possible event interleavings caused by concurrency and failures. Using a multicast primitive that provides ordering and reliability guarantees, reduces the possible event histories, making correctness arguments simpler.

Synchronous primitives, such as synchronous send and remote rendezvous, have been popular in the functional language community, because of their simple ordering semantics. But with synchronous communication it is difficult to exploit the full bandwidth of fast computer networks [Birman and van Renesse 1992, Knabe 1991] because each communication necessarily involves one or more round-trip packet exchanges over the network. When multicast is considered, these costs become intolerable. Asynchronous communication permits pipelining for good performance, and uses new ordering properties to substitute for those provided by synchronous primitives.

Distributed ML [Krumvieda 1991] presents flexible, asynchronous group communication facilities using the higher-order concurrency primitives of Concurrent ML [Reppy 1991].

I. INTRODUCTION

Standard ML (SML) [Milner et al. 1989] is a modern language that supports the construction of correct programs. For instance, SML provides first-class functions, strong static typing, polymorphism, and exception handling. Several features of the language make it especially appealing to the distributed programmer:

• Most objects in SML are immutable values; unlike C functions, SML applications cannot directly modify complex structures by assigning to their internal fields. Because an immutable
function parameter can be passed either by value or by reference (both methods produce the
same results), a distributed programmer need not worry about semantic differences between
local and remote evaluation.

- Many distributed applications evolve from separately running programs that use the network
to share common data. These so-called multi-applications [Auerbach et al. 1991] and their
interfaces must be designed well if they are to be combined effectively. SML's module
system—which allows module interfaces to be type-checked and parameterized in terms of
each other—provides a good framework for designing multi-applications.

- Standard ML has a particularly efficient implementation, Standard ML of New Jersey
(Appel and MacQueen 1991), whose code seems to run within a factor of two of optimized
C code. Since software costs constitute a significant fraction of the total costs of distributed
communication, the low-level support system of a distributed language must be efficient.

- Any general purpose distributed programming system must provide support for node-level
(lightweight) concurrency. Concurrent ML (CML) [Reppy 1991], an extension of SML,
adds high-level support for single-environment concurrent programming. CML is especially
appealing because it provides excellent performance.

At least two research groups have implemented distributed computing extensions to SML
[Knabe 1991, Matthews 1991]. Both groups have chosen to implement synchronous com-
munication, where data sources (senders) rendezvous with data sinks (receivers) to transfer data.
Synchronization between senders and receivers ensures the stability and ordering of message de-
ivery, thereby aiding program correctness. Unfortunately, synchronous communication makes it
difficult to exploit the full bandwidth of fast computer networks [Birman and van Renesse 1992].
In Distributed ML (DML) [Krumvieda 1991], senders do not wait for receivers to rendezvous
with them, but instead use asynchronous communication. We will argue that asynchronous com-
munication primitives are better than their synchronous counterparts for distributed programming
languages and explain how the stability and ordering problems are addressed.

Although point-to-point (single sender, single reciever) communication in a distributed system
is important, distributed applications frequently employ one-to-many (multicast) communication,
e.g., to efficiently disperse data and maintain replicated data sets. Multicast can be more difficult to
use than point-to-point communication because of the larger number of possible event orderings.
But by adding some reliability and ordering properties [Birman et al. 1991b], the possible orderings
can be substantially reduced and correctness arguments simplified. We will argue for reliable,
ordered multicast as a primitive in a distributed programming language.
II. DISTRIBUTED ML

A DML program comprises a set of nodes which contain multiple pre-emptive threads sharing access to a heap. Nodes may fail by crashing.

Nodes communicate with each other via port groups, collections of src.ports (source ports) and dest.ports (destination ports) used to multicast data (see Figure 1). A port group serves the same purpose in DML that a channel does in CML: it transmits typed data between threads. Conceptually, data which are placed on a group's src.port are transported to each of the group's dest.ports.

![Fig. 1. Conceptual view of a port group](image)

Groups have meta.src.ports and meta.dest.ports which convey information about port group membership. For instance, the information that a certain src.port has failed appears on the group's meta.dest.port. Meta ports are examined in detail in a companion paper [Krumvieda 1992].

An abbreviated signature for ports and port groups appears in Figure 2. The portTransmit function is used to asynchronously transmit a datum through a src.port. It returns a CML.event value, which can be used to obtain a message delivery guarantee (see Section V). Data sent through port groups are sent asynchronously, and destination ports act as data buffers.

Port groups with more than one src.port must interleave data that are multicast to their dest.ports. The programmer specifies one of the orderings defined in the ORDERINGS signature to mkGrp (discussed in Section IV) when creating the group.

The portReceive function is used to receive a value from a destination port. It returns an OrdEvt.ord.event value that encodes multicast ordering information. How ordered events differ from CML events is discussed in Section VI.
III. MULTICASTS

Just as point-to-point message primitives benefit from properties such as reliability (e.g., retransmitting messages lost by the network) and FIFO ordering, multicast communication is easier to use and reason about if it possesses failure and ordering properties [Birman et al. 1991a]. Two of these properties are listed below.

- **Multicast atomicity**: a multicast should be delivered to either all or none of its destinations. A multicast might not be delivered if its sender crashes part way through its transmission.

- **Multicast ordering**: a multicast may be ordered relative to other multicasts in the system. The three properties below generalize the FIFO property common in point-to-point commu-
- **FIFO ordering** requires multicasts to be ordered pair-wise between the sender and each receiver.

- **causal ordering** preserves potential causality between sends and receives, and is discussed in detail in Section IV.

- **total ordering** ensures that all multicasts are delivered in the same order at all destinations. It enforces an ordering on concurrent multicasts initiated by different senders, and also observes the causal ordering property.

Correctly implementing ordered, reliable multicast out of simple point-to-point primitives is non-trivial (whether based on synchronous or asynchronous point-to-point primitives), and deriving such protocols has been an active research topic for the last decade. Moreover, many networks, including Ethernet, support **hardware multicast** which transmits a multicast as quickly as a single point-to-point message. Once a multicast has been translated into numerous point-to-point messages, it is extremely difficult—or at least very expensive—to reconstitute the multicast destination sets. Therefore, distributed languages that wish to support multicast applications should provide a multicast primitive. Although one could imagine a system that provided a synchronous multicast—one in which a sender synchronized with all receivers—such a primitive would be extraordinarily expensive, especially if it could be used in selective communication [Knabe 1991]. DML provides an asynchronous multicast.

### IV. MULTICAST ORDERING

An important property of synchronous communication is that any operations executed by the sender upon return from the synchronous send are known to occur after the receipt of the message at the destination. That is, synchronous send operations force particular orderings on the events that follow a communication at both sender and receiver. Often these orderings are too strict and can be substituted with cheaper orderings. DML port groups support three kinds of ordering: FIFO, causal, and total. Causal ordering is the least widely understood and the most interesting from theoretical and performance aspects.

#### A Causal Ordering

Causal ordering is based on the potential causality (or “happens before”) relation [Lamport 1978]. The **causal ordering** method ensures that if the sending of \( m' \) causally follows the sending of
m, then each destination that receives both messages delivers m before m' [Birman et al. 1991b]. Consider the example in Figure 3 concerning three threads, P, Q and R. P sends message $m_1$ to R and then sends $m_2$ to R. Upon receiving $m_2$, Q sends $m_3$ to R. It is probably intended that R receive $m_1$ before $m_3$; at first glance, it seems that synchronous message passing would be necessary to enforce this behavior. However, it is much more efficient to use causally ordered asynchronous messages. Causal ordering forces R to receive $m_1$ before $m_3$ without requiring an explicit acknowledge message. Causal ordering can be used in many instances where a synchronous protocol might at first appear necessary.

Using asynchronous send operations, we can build a synchronous abstraction from a pair of asynchronous message operations and use it when synchronous properties are desired. In DML, this abstraction would have the same status as the built-in asynchronous operations. Of course, one could argue that asynchronous operations can be implemented using synchronous operations and extra threads. But this implementation incurs the extra message transmission costs we wish to avoid in the asynchronous case [Birman and van Renesse 1992].

Fig. 3. Causal ordering
signature ORDERINGS =
  sig
    type causal_dom
    type total_dom
    datatype ordering = FIFO
    | CAUSAL of causal_dom
    | TOTAL of total_dom

  val causalO: causal_dom
  val mkTotDom: causal_dom -> total.dom
  ...
end;

Fig. 4. Ordering signature

B Causal Completeness

DML’s causal ordering implies a causal completeness property. Informally, this property guarantees that there are no gaps in the middle of a process’s causal message history due to crashes, although the history may be truncated by a crash. More precisely, causal completeness states that for any message, m, delivered to process P:

- all m’s causal predecessors that are destined for P will have been delivered to P, and
- all m’s causal predecessors not destined for P will eventually be delivered to their surviving destinations, despite subsequent node crashes.

There are a range of implementation techniques that satisfy this property.

C Ordering Declarations

In DML, orderings are associated with port groups. Recall that the mkGrp function from Figure 2 had an ordering argument. The signature for orderings is in Figure 4.

Elements (domains) of type causal.dom and total.dom may be used to enforce ordering schema among multiple port groups. For instance, if two port groups were declared to be CAUSAL with respect to the same causal.dom, data multicast through either the group is ordered relative to data multicast in both groups.

V. MESSAGE STABILITY

One important property of synchronous communication is that when a synchronous send operation returns, its message has been reliably received by the destination node(s). In particular, the
subsequent failure of the sender cannot prevent delivery of the message to the receiver. Message
stability can be expensive, because a sender must wait after transmitting a message until an
acknowledgement is received from a receiver. As computer networks and I/O hardware become
faster, they provide larger message throughput but spend relatively more time putting messages on
the wire. A sender and receiver cannot use this higher bandwidth effectively if every communication
must incur the round-trip packet latency of the network.

In practice, message stability need be achieved only at particular points in a stream of remote
sends (e.g., before dispensing cash from an automatic teller, before confirming a reservation to a
ticketing agent, or before closing a file). DML's portTransmit operation does not guarantee
message stability, but returns a CML event value, a token which can be redeemed for a guarantee
that the message has been received at its destinations.

In many other cases the multicast atomicity property suffices. The portTransmit operation
implements multicast atomicity when sending to multiple destinations.

VI. ORDERED EVENTS

One of the more remarkable features of Concurrent ML is its support of first-class synchronous
operations [Reppy 1991]. Many concurrent operations (e.g., timing out, message transfer, detection
of thread termination) block, and blocking operations are not easily embedded in new synchronous
operations. In CML, concurrent operations return event values1 instead of blocking, which permits
new synchronous constructs to be constructed easily and elegantly.

Unfortunately, CML events are unordered and therefore not appropriate for some DML opera-
tions. For instance, consider the CML function

\[
\text{select: } 'a \text{ event list} \rightarrow 'a
\]

which synchronizes upon exactly one element of its parameter list. If more than one element is
ready for synchronization, the function chooses one of them.

Consider using CML's select on a list of two receive events ordered by the causal ordering
property. If both receive events were available when the select was executed, it would be
inappropriate to choose nondeterministically between them, and instead, the prior event should be
selected. One could pass all ordered receives through a thread that released them one at a time, but
doing so imposes a total order when only a partial order suffices, thereby restricting concurrency
and, perhaps, leading to deadlock. Indeed, the Isis system [Birman et al. 1991a], from which these
ordering and multicast properties are derived, constrains each node to receive a totally ordered
message stream.

1 An event should be thought of as the potential to block. In particular, it is not a future, and functions which return
event values need not initiate a blocking operation.
signature ORD_EVENT =
  sig
    structure CML: CONCUR_ML
    type 'a ord_event
    val resolve: 'a ord_event list -> 'a ord_event
    val unord: 'a ord_event -> 'a CML.event
    val sync: 'a ord_event -> 'a
    val wrap: 'a ord_event * ('a -> 'b) -> 'b ord_event
    ...
  end

Fig. 5. Ordered event signature

In contrast, DML provides ordered events that, while similar to CML events, behave as elements of a partial order. Part of the ORD_EVENT signature is in Figure 5. The resolve operator selects between ordered events in a manner consistent with the underlying partial order. The unord function strips ordered events of their ordering information, and the sync and wrap operators operate like their CML counterparts. Ordered events allow DML to inherit the nice features of CML while preserving the information required to support ordered asynchrony in a pre-emptive environment.

VII. EXAMPLE: REPLICATED PROCESSING

A more substantial example will illustrate both the ordering and stability issues. Suppose we are given a distributed program that communicates using only point-to-point channels (see Figure 6); consider how we can make it tolerate single crash failures by replicating each node [Alsberg and Day 1976]. A sketch of a non-fault-tolerant implementation of the POINT-TO-POINT signature appears in Figure 7.

In the fault tolerant version we replace each node \( P \) by a pair \( (P, P') \) consisting of a primary node and a backup which will take over the role of the primary should it fail. The backup maintains its internal state equivalent to the primary’s, and, upon the primary’s failure, produces a sequence of output messages consistent with the state of the primary at the instant it crashed. To achieve this, we arrange for the backup to observe exactly the same input messages as the primary, in the same order. Where the execution of the primary is nondeterministic (e.g., because of pre-emptively scheduled threads), we must ensure the backup takes the same nondeterministic decisions. We will concentrate on the ordering and atomicity properties of internode communication, ignoring other
signature POINT_TO_POINT =

sig
signature PortGroup: PORT_GROUP

  type 'a remote_channel
  (* Only one dest_port is permitted per channel. *)

  val mkChannel: unit -> 'a remote_channel
  val mkSrc: 'a remote_channel -> 'a PortGroup.src_port
  val mkDest: 'a remote_channel -> 'a PortGroup.dest_port
  val portSend: 'a PortGroup.src_port * 'a -> unit
  val portAccept: 'a PortGroup.dest_port -> 'a

end

Fig. 6. Signature for simple point-to-point remote communication

functor SimpleP2P(structure PortGroup: PORT_GROUP): POINT_TO_POINT =
struct
  structure PortGroup = Portgroup

  local open PortGroup PortGroup.Ports
  in
    type 'a remote_channel = 'a port_group,

    fun mkChannel () = mkGrp Ord.FIFO;
    fun mkSrc g = #1 (PortGroup.mkSrc (g, ()));
    fun mkDest g = #1 (PortGroup.mkDest (g, ()));
      (* In practice there would be code to ensure only one
         dest_port per channel. *)
    fun portSend (src, data) = (CML.sync (portTransmit (src, data)); ());
    fun portAccept dest = OrdEvt.sync (portReceive dest);
  end;
end

Fig. 7. Non-fault-tolerant implementation of point-to-point channels
signature PROCESS_PAIR = 
Sig
    val primary: bool ref
    type schedule (' Information about nondeterministic decisions made
        by primary *)
    val backup_chan: schedule port_group
end

functor ResilientP2P(structure PortGroup: PORT GROUP
    and ProcessPair: PROCESS_PAIR): POINT_TO_POINT =
struct
    structure PortGroup = Portgroup

    local open PortGroup PortGroup.Ports
    in
        type 'a remote_channel = 'a port_group,

        fun mkChannel () = mkGrp Ord.TOTAL;
        fun mkSrc g = #1 (PortGroup.mkSrc (g, ()))
        fun mkDest g = #1 (PortGroup.mkDest (g, ()))
        (* In practice there would be code to ensure only one dest_port
            at each of the primary and backup. *)

        fun portSend (src, data) =
            if ProcessPair.primary then
                (CML.sync (portTransmit (src, data)); ())
            else ()

        fun portAccept (src) =
            OrdEvent.sync (portReceive src);
        end;
    end
end

Fig. 8. Fault tolerant implementation of point-to-point channel.

issues including how the backup would use a meta_dest_port to notice the failure of the primary
and how to handle nondeterminism due to thread scheduling and external I/O [Borg et al. 1989,
Birman et al. 1991a].

We represent each point-to-point communication channel in the original program by a port
 group containing two dest_port's, one owned by each of the primary and backup nodes (see
Figure 8). The backup reads these messages and performs the same actions as the primary, except
that output messages from the backup are suppressed. When the primary fails, the backup assumes
the primary's role, setting the variable ProcessPair.primary to true. In addition there is a
point-to-point port group (ProcessPair.backup_chan) from the primary to the backup. The
primary uses this connection to send messages to the backup that resolve any nondeterminism, for
A point-to-point message, $m$, from node $P$ to $Q$ in the original program is transformed into a multicast from $P$ to $Q$ and its backup $Q'$ (see Figure 9). Using a totally ordered multicast, $Q$ and $Q'$ will receive this message in the same order, relative to other multicasts.

We must also ensure that the scheduling message, $b$, sent by $Q$ over the backup channel is delivered at $Q'$ after the causally preceding multicast, $m$. If synchronous, non-causal communication is used, $P$ must wait when it-transmits $m$ until both $Q$ and $Q'$ have received and acknowledged the message. If causally ordered communication is used throughout, $P$ can initiate the multicast and immediately continue processing while the message is still being delivered. The causal ordering protocol includes information in message $b$ identifying $m$ as a causally preceding message that must be delivered first. With synchronous, non-causal communication, an extra message delay is inserted into the critical path of the program.

This performance penalty is more severe than at first appears. To increase the independence of failures we might locate the primary and backup of a pair in physically separate locations. To improve performance in the normal case (no failures) we might locate all the primaries near each other (perhaps on the same computer). Now the synchronous approach performs much more slowly than the asynchronous causal approach. In the synchronous case, all communication will
occur at the speed dictated by the primary-backup connections. In the asynchronous approach, communication among the primaries can proceed at close to the speed attainable in the original non-fault-tolerant program. Communication between primaries and backups can proceed slower, in the background. Because messages are asynchronous, multiple primary-backup messages can be combined automatically into a smaller number of large network packets which will make much more efficient use of the network. We are not limited by network latency (round trip packet times), but bandwidth. Bandwidth scales much better than latency on almost all network technologies.

But what happens if a primary fails before its backup node has received all of the messages destined for it? The causal completeness property ensures that for any message received by another node, the causally preceding messages destined for the backup will be delivered to it. Thus, if any "evidence" of an action taken by the primary has been observed by another node in the program, the backup will receive any prior messages from the primary. Conversely, if no evidence of the final few actions of the failed primary is visible, then we can present the illusion that those actions never took place. There remains the possibility that the backup will send out some messages that duplicate the last few messages sent by the primary. These duplicates can be detected easily at the destinations using sequence numbers.

We see that asynchronous communication would not be feasible in this example without the causal ordering and completeness properties, which ensure that asynchronous messages are delivered in the correct order even when they are delayed and nodes crash, and the multicast atomicity property, which ensures that either all destinations receive the multicast, or none of them do.

VIII. CONCLUSION

We have argued that asynchronous communication, coupled with appropriate ordering and failure properties, permits higher network performance than synchronous communication. As networks become faster relative to processors, asynchronous communication will be essential. Additionally, we have argued for multicast as a communication primitive in support of fault tolerant and parallel programming. DML's ordering properties, such as total and causal orderings, are essential to both multicasts and asynchronous communication. They reduce the number of possible event histories, without compromising performance.
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Summaries of Poster Session Presentations
A Verification Environment for ML Programs

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Abstract

A verification environment for Standard ML programs is described. The system is constructed within the
Isabelle theorem prover, directly from the operational semantics definition of $F$, the functional subset
of the Core of SML. Both the static and the dynamic semantics of $F$ are incorporated. Simple proof
procedures can prove $F$ programs correct by inferring the result of evaluation or elaboration demanded
by the Definition. The work will benefit those trying to understand the Definition of SML, and experts
who wish to explore possible modifications and extensions.

1 Introduction

In this paper we report on progress towards the construction of a verification environment for reasoning
about ML programs. The system we describe captures the formal operational semantics definition of
Standard ML [1] within the theorem prover Isabelle. Our work has arisen from the desire to explore
the role of formal semantics in the verification process, especially in the context of critical software, in
which a single error could have disastrous consequences. It complements work described in [2], where
we explored the role of denotational semantics in the verification process.

ML has evolved from its early days as a special purpose language for theorem provers [3, 4] to the
point where it is a serious programming language, expressive enough for many real applications, and
with numerous desirable features (such as strong typing, exception handling and modules) which modern
software engineering and critical software development require. Although, to our knowledge, ML has not
yet been used in critical software projects, we believe that the techniques described below are applicable
to any language built on rigorous principles. They will also be applicable to the problem of automated
reasoning about programs in process algebras such as CCS [5] and CSP [6].

The rest of this paper is structured as follows. In Section 2 we describe the structure of the Definition
of SML. In Section 3 we give a brief overview of Isabelle. Section 4 describes how our system is
constructed. Finally, Section 5 discusses the results and give suggestions for further work.

2 The Definition of SML

The rigorous basis for Standard ML is formally given by the Definition [1], and its companion Com-
mentary [7]. To quote from [1]:

"This document is a formal description of both the grammar and the meaning of a lan-
guage which is both designed for large projects and widely used. ... At a time when it is
increasingly understood that programs must withstand rigorous analysis, particularly for
systems where safety is critical, a rigorous language presentation is important even for ne-
gotiators and contractors, for a robust program written in an insecure language is like a
house built upon sand."

The Definition uses operational semantics [8] to describe the meaning of phrases in the language. This
method of language definition is a useful guide for the implementer of an interpreter or compiler for
the language. It is especially useful for describing the semantics of concurrent languages, where the
denotational semantics may be technically difficult and less easy to mechanise.
The Definition first presents the syntax for SML (both for the Core language and the Modules System), introducing various identifier and phrase classes. The static semantics (elaboration) is then given in detail. This involves a rich set of simple and compound semantic objects (such as environments, types etc). Elaboration of a phrase is expressed by a sequent of the form

\[ E \vdash \text{phrase} \Rightarrow \text{result} \]

where typically E is an environment, and the result may be a type or an environment. The 102 inference rules capture all the possible inferences among these sequents.

The dynamic semantics is then given a similar, but quite separate, treatment. The fact that evaluation and elaboration can be dealt with independently is an important aspect of SML. Often the same terminology gets used in both the static and dynamic semantics but has a different meaning in each case, such as "variable environment". Static and dynamic semantics meet at the level of programs, where the evaluation of a program is only carried out if it elaborates successfully.

Standard ML has a number of phrase classes which are derived forms. For example, the program phrase case exp of match is defined to be the more primitive language expression (fn match) (exp). Other examples of derived forms are if, andalso and orelse, as well as lists and tuples. Inference rules only need to be given for the phrases in the so-called bare language.

Because of their formal nature, and the size of the language, the Definition and Commentary are not light reading, and are aimed at implementers and ML experts more than the general reader. The Definition allows one in principle to explore language semantics, but detailed proofs done on paper using all the 196 inference rules are far too laborious: we believe that machine support is essential to be able to do this.

In constructing a system for reasoning about SML programs, our aim has been to capture the definition of SML in as natural a way as possible in a powerful proof assistant. One might consider using Prolog for this purpose [9]. However, we have chosen to use the theorem prover Isabelle, which has a number of advantages over Prolog: it allows the use of concrete syntax; theories can easily be related to familiar logics (such as First Order Logic and Higher Order Logic); and one has much finer control of proofs steps and search. We believe, therefore, that Isabelle is an ideal tool for reasoning about operational semantics.

3 Isabelle

The theorem prover Isabelle (itself constructed using SML!) has been under development by Larry Paulson at the University of Cambridge since 1986 [10, 11, 12]. Isabelle is a generic theorem prover, with an expressive meta-logic, in which the inference rules and axioms of object logics can be formulated. Isabelle emphasises the natural style of reasoning, and thrives on inference rules such as those of operational semantics. It is this fact which has been exploited in our work.

Isabelle carries out goal-directed proofs. A proof state consists of a goal, along with a number of subgoals whose validity establishes that of the goal. Proving a goal involves reaching a proof state with no subgoals, by means of the application of tactics, which transform proof states to new proof states. In Isabelle a tactic may fail, or return one or more (and possibly even an infinite number) of new proof states.

The most important tactic is resolution: resolve_tac thms i tries each theorem (object logic rule) in the list thms against subgoal i of the proof state. For a given rule, say

\[ \{ B_1, \ldots, B_k \} \Rightarrow B \]

resolution can form the next state by unifying the conclusion with the subgoal, replacing it by the instantiated premises. (Note that unification in Isabelle is full higher-order unification [13]). Thus if the subgoal is

\[ \{ A_1, \ldots, A_n \} \Rightarrow A \]

and A can unify with B, resolution will produce the following new subgoals:

\[ \{ \overline{A_1}, \ldots, \overline{A_n} \} \Rightarrow \overline{B_1} \]

\[ \ldots \]

\[ \{ \overline{A_1}, \ldots, \overline{A_n} \} \Rightarrow \overline{B_k} \]
in which the overbars denote the resulting formulae after instantiations have been made.

Isabelle also has a number of *tacticals*, used for building new tactics from basic tactics, for example:

- tac1 THEN tac2 (sequencing)
- tac1 ORELSE tac2 (choice)
- \textbf{DEPTHFIRST} pred tac (search)

The tactic \textbf{DEPTHFIRST} \texttt{pred tac} performs a depth-first search for a proof-state satisfying \texttt{pred}. Usually \texttt{pred} is taken to be "no subgoals", so that the tactic will search for a proof of the original goal.

Isabelle also has answer extraction available, via so-called *scheme variables*. These variables can be part of a goal; as tactics are applied the scheme variables may be instantiated during the proof.

4 Description of the System

At present, we have built a system for a non-trivial subset \( \mathcal{F} \) of Standard ML — essentially the pure functional (side-effect free) subset of the Core Language, including pattern matching, functions as first-class objects and recursion. Therefore, we have excluded imperative features such as reference variables, assignments, and exceptions, as well as the modules system. However, what remains is still an extremely rich language.

The syntax, semantic objects and inference rules of \( \mathcal{F} \) are constructed as a new theory which is a (typed) Isabelle object logic \( \mathcal{L} \). The types of the logic are expressions, declarations as well as objects such as values, environments, identifiers etc.

4.1 Syntax

Isabelle allows the user considerable freedom in the choice of concrete syntax. It is a great aid to understanding to be able to use a nice concrete syntax which is also strictly maintained during a proof. Our aim has been to use wherever possible the precise syntax of SML. For example, functions are given in SML by:

\[
\text{exp ::= ...}
\]
\[
\text{fn match}
\]
\[
\text{match ::= pat => exp (| match)}
\]

which is easily expressed as the following Isabelle syntax declaration:

\[
\text{Delimfix (\text{"fn \_\"}, Match --> Exp, ...)},
\]
\[
\text{Delimfix (\"\_ => \_\", \[Pat, Exp\] --> Match, ... )},
\]
\[
\text{Delimfix (\"\_ => \_ | \_\",}
\]
\[
\text{[Pat, Exp, Match] --> Match, ...),}
\]

Note that Match and Exp appear as new types in the logic; thus Isabelle's type-checking will catch syntax errors.

However, variables (and also value constructors and special constants) do need to be explicitly marked, for example:

\[\text{fn \{var x, var y, \ldots\} => var x + var y}\]

Note that we are at liberty to overload notation: for example, we can use curly brackets for record expressions, record patterns and record values, just as the Definition requires. The same holds for lists and pairs. Isabelle can tell what is meant from the context.

Another of Isabelle's strengths is the ability to include derived forms by means of parse translations (which take a simple concrete syntax phrase such as \texttt{case exp of match} into the internal representation of the appropriate bare language phrase), and print translations (which reverse the process). Once these translations have been provided, we can use the derived forms freely in goals, and they will be correctly dealt with.

\textsuperscript{1} Actually, \( \mathcal{L} \) is an extension of First Order Logic.
4.2 Semantic Objects

In the spirit of [1], elaboration (static semantics) and evaluation (dynamic semantics) are treated separately; this is reflected in the design of the system. Thus, separate Isabelle theories are maintained: one for elaboration and one for evaluation. They have in common the syntax of $\mathcal{F}$ itself.

The various semantic objects, such as values, environments etc are themselves given an appropriate concrete syntax, and their properties described by means of inference rules within the logic $\mathcal{L}$ for the various operations defined on these domains. For example, variable environments are of the form

$$\{ (x_1, v_1), \ldots, (x_n, v_n) \}$$

for evaluation, and of the form

$$\{ (x_1 : t_1), \ldots, (x_n : t_n) \}$$

for elaboration.

Typical inference rules are the following, which describe the operation of looking up a value in a variable environment:

```isabelle
val lookup =
  ["lookup1_rule",
   " lookup (x, \{ (x, v) \}, v)",
   "lookup2_rule",
   " lookup (x, \{ (x, v), bindseq \}, v)",
   "lookup3_rule",
   " lookup (x, \{ bindseq \}, v) =>
     lookup (x, \{ (y, w), bindseq \}, v) " ];
```

4.3 Static Semantics

For the most part, the inference rules capturing the static semantics of $\mathcal{F}$ are easily expressed in Isabelle. For example, here is the Rule 6 for the elaboration of the atomic expression let dec in exp end. This is expressed as the following Isabelle fragment

```isabelle
("Lettyrule", (* Rule 6 *)
  "[| C \|- dec -> E ; combine (C,E,C') ; C' |- exp -> t |
  ==> C \|- let dec in exp end -> t ")
```

where combine is the operation which combines contexts together. Note the way that sequents are written in these rules.

In the static semantics, the trickiest aspect to model is polymorphism. We need to set up a number of inference rules to implement the considerable machinery of type schemes, type instances and closures in order allow sound polymorphic typing.

4.4 Dynamic Semantics

Once again, it is straightforward to express inference rules. Consider, for example, the Rule 108, for the evaluation of the above expression:

```isabelle
("Let_rule", (* Rule 108 *)
  "[| E \|- dec -> E' ; combine (E,E',E''); E'' |- exp -> v |
  ==> E \|- let dec in exp end -> v ")
```

The dynamic semantics needs to have function closures, in order to have correct call-time environments. Recursion demands the unfolding operation on environments. These have been incorporated in the system.

4.5 Proof Procedures

The proof procedures for reasoning about programs are quite simple Isabelle tactics which capture the way one would construct an inference tree [7]. Starting from the initial goal (the root of the tree), we
keep resolving with the appropriate inference rules, simplifying environments as we go, until all language phrases have disappeared. We then simplify using the inference rules for semantic objects until we reach the leaves of the tree. Isabelle keeps track of the instantiations made as we go. Thus we can prove $\mathcal{F}$ programs correct by inferring the result of evaluation or elaboration demanded by the Definition. Proofs of ML programs involving pattern matching may require backtracking search. To accommodate this, our proof procedures use the depth-first search tactical.

Polymorphic type inference, as one would expect, requires special handling. However, the actual work of inferring the most general type of an expression is aided by Isabelle’s scheme variables, which can then be instantiated to type variables.

5 Conclusions and Suggestions for Future Work

The system is easy to modify, and is quite efficient, because it exploits Isabelle’s liking for inference rules, and keeps costly rewriting to a minimum.

We believe that the present work will benefit a number of people. It should be of help to beginners trying to understand the Definition of SML, as well as implementers. It can assist experts who wish to explore possible design changes and extensions to the language, by aiding reasoning about how the various phrases will interact. We also believe that the system will benefit those interested in program verification, equivalences and transformations.

The user interface is still under development. Ideally, it should offer the user a range of choices. A minimal interface would be to mimic that of an ML interpreter, presenting only the value and type of the most recently declared variable(s). The maximal interface would be to describe fully the proof of each sequent, showing also the context before and after the evaluation or elaboration. This information can quickly become too much to cope with.

In future work, we would like to extend the system to allow for a larger subset of SML. Exceptions and imperative features will require proper handling of the state and exception conventions [1, 7], while the inclusion of the modules system would allow experiments with subtle aspects of signature matching and elaboration of functors and signature expressions.

Further work needs to be done on general proofs of correctness of algorithms written in SML, for which we need to have an expressive specification language. Since our system is built on Isabelle’s First Order Logic, we already have the basis of such a language, while our system could easily be built on Isabelle’s Higher Order Logic, if required. Such specification constructs are under investigation as part of the Extended ML language of Sannella and Tarlecki [14], and in current work of Gene Rollins, Jeannette Wing, and Amy Moormann Zarembski at CMU on Larch/ML.

Another important line of investigation is that of reasoning about program equivalences and transformations [15, 16]. The operational semantics rules which define SML will need to be strengthened to allow such reasoning.

We are also planning to apply our methods to concurrency, where operational semantics is frequently used to give the meaning of language constructs: possibilities include CML [17], CCS [5] and the tasking model of Ada.

6 Acknowledgment

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Expressing Fault-Tolerant and Consistency-Preserving Programs in Distributed ML

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ABSTRACT

Many of the programming problems particular to distributed environments involve violations of consistency. For example, some applications require that replicated data copies be kept identical. If explicit failure information is provided by the programming system, failure notifications must be ordered so that distributed components maintain a consistent view of system membership. We are currently adding features to the Standard ML (SML) programming language that help distributed programmers preserve consistency. Our system, called Distributed ML (DML), introduces ports and first-class port groups as a method of abstracting asynchronous multicasts. DML provides meta ports, multicast orderings, and ordered events to help programmers reason about the dynamic distributed environment. This paper contains three examples that illustrate DML's syntax and operational semantics.

I. CONSISTENCY

There are at least two reasons that programmers write distributed programs. First, many applications require parallelism. For instance, distributed databases and electronic mailers are inherently distributed; other applications need real concurrency to execute quickly. Second, some applications must behave correctly despite component failures. Running a single program on multiple, independently-failing computers is an appealing way to achieve resiliency. File servers and air traffic controlling programs are examples of applications requiring high availability.

Unfortunately, naive attempts to implement fault-tolerance and concurrency with distributed systems can lead to violations of consistency [Birman 1991]. There are numerous consistency properties, and they vary among applications. For instance, some applications maintain replicated data sets which must remain consistent with each other. Data consistency is threatened if replicas receive concurrent updates in different orders. Also, many applications divide work among active processes and therefore require a consistent view of outstanding tasks and available processors. Component failures may lead to inconsistent system "views" of active processes.

In a companion paper [Cooper and Krumvieda 1992], we discussed the Distributed ML (DML) programming language and justified its asynchronous multicast primitives. Port groups, multicast orderings, and ordered events are several of the tools that DML programmers can use to maintain consistency in a distributed environment. In this paper, we present three DML programming examples and argue that explicit failure information through meta ports can be used to maintain consistent views of system membership.

II. EXAMPLE 1: RPC MULTICAST

The client/server model is a common paradigm in fault-tolerant distributed programming. In this model, a group of server programs control a resource (e.g., a database or a speedy processor) while client programs...
communicate with the servers to access the resource. Often, the servers are replicated to provide fault
tolerance. Here, communication is two-way; that is, clients initiate communications with a server and
servers send reply messages to waiting clients. If a server group has only one member, this communication
style is called RPC, or Remote Procedure Call. If there are several servers, the style is RPC Multicast; often,
distributed toolkits (such as Isis [Birman and Joseph 1987]) directly support RPC Multicast.

One possible DML implementation of RPC Multicast would require a single port group shared by both
clients and servers; clients would ignore messages sent by other clients. However, this solution requires
each client to have a separate dest_port and would be too inefficient when clients greatly outnumber
servers (which is typical).

Instead, we define a type ('a, 'b) rpc_group. Clients send requests of type 'a to servers, while
servers send replies of type 'b to clients.

```haskell
type 'a rpc_reply = 'a * port_id
type ('a, 'b) rpc_request = 'a * ('b rpc_reply, unit) port_group
type ('a, 'b) rpc_group = ('a, 'b) rpc_request, unit) port_group
```

Data sent through an rpc_group are rpc_requests, pairs which contain a request and a port group
for replies. Replies are pairs containing reply data and a port_id identifying the replying server. All meta
data in this example has type unit.

The function request, used by clients to initiate RPC multicasts, has type

```haskell
val request: ('a, 'b) rpc_request, unit gview event) src_port * 'a ->
'b rpc_reply dest_port * unit gview event
```

The first coordinate of the pair returned by request is a destination port used to queue server replies;
each reply contains data of type 'b and a port_id of the dest_port from which the request was received.
The second half of the pair is a unit gview event which, when CML_sync is applied to it, yields the
membership of the server group when the original message was delivered\(^1\). This information is enough to
collect replies in a number of different ways, one of which we will examine in section IV.

The server uses the function service to reply to client queries.

```haskell
val service: ('a, 'b) rpc_request dest_port * ('a -> 'b) -> unit
```

This function is used to register a "handler" function with a destination port. After a service call is
evaluated, incoming requests are processed automatically; replies are calculated by feeding requests to the
registered function.

The implementations of request and service are shown in Figure 1. At first glance, they seem to
be extremely inefficient because the number of port groups required is linear in the number of requests
(assuming a bounded number of servers). However, port groups with only one destination port can be
implemented with point-to-point communication and are therefore cheap.

RPC Multicast clients receive a stream of reply messages and, to avoid blocking for messages that may
never arrive, they must monitor the servers for failure.

### III. EXPLICIT FAILURE INFORMATION

DML is intended to be used in asynchronous distributed systems, that is, systems in which communication
times are unbounded\(^2\). Unfortunately, many important problems that involve consistency among processes—

\(^1\)The unit in unit gview event is the group's meta data type and will be discussed in section III.

\(^2\)The word "asynchronous" is overloaded: asynchronous systems should not be confused with asynchronous communication. Asynchronous communication (in which data senders do not rendezvous with data receivers) can occur in synchronous systems and vice versa.
fun request (sp, msg) = let
  val rg = mkGrp FIFO;
  (*1 (mkDest (rg, ())), portTransmit (sp, (msg, rg)))
end;

fun service (dp, fcn) = (wrap (unord (portReceive dp),
  fn (msg,rg) => let val sp = (*1 (mkSrc (rg, ())));
    (*1 portTransmit (sp, (fcn msg, portID dp))); ()
  end);
  service (dp, fcn));

Fig. 1. Implementations of request and service

such as agreement on system membership and consensus on the value of a variable—cannot be solved in failure-prone asynchronous systems [Fischer et al. 1985].

The DML runtime system circumvents this impossibility result by providing a (scalable) system-wide membership service [Ricciardi and Birman 1991]. The membership service maintains the set of active DML nodes and can add and remove nodes dynamically. In particular, the service may delete nodes that appear to have failed (perhaps because they didn’t acknowledge a series of messages), even if they may still be working properly.

DML port groups have meta ports that the membership service uses to inform programs of changes to the system membership. Consider a portion of the signature PORT-GROUP (more of which is defined in [Cooper and Krumvieda 1992]):

signature PORT-GROUP =
sig
  structure Ports: PORTS

  type 'a gview
  type 'a monitor_event = SRC_CREATE of 'a * Ports.port_id
  | SRC_FAIL of Ports.port_id
  | DEST_CREATE of 'a * Ports.port_id
  | DEST_FAIL of Ports.port_id

  val getView: ('a, 'b) port group -> 'b gview
  val portMonitor: 'a gview -> ('a monitor_event * 'a gview) Ports.dest_port

end;

A value of type gview contains port group membership information, and a monitor_event describes a specific event that triggered a change in the membership of some port group. All events provide the port_id of the created or failed port; create events also carry information provided when the port was created. The getView function uses the membership service, if necessary, to take a “snapshot” of its port group argument. The portMonitor function examines its gview argument, adds a meta dest_port to the appropriate group, and enqueues, at the beginning of the new dest_port, all changes (monitor_events) that the group underwent since the gview was constructed. Conceptually, the dest_port returned by the portMonitor functions is a stream of all membership changes—past, present, and future—to the port group described by its gview argument since the view was created.
fun procReply (n, (dp, ge)) = let

  fun statel got =
    if length got = n then []
    else sync (choose (wrap (unord (portReceive dp),
                                fn (x, id) => x :: (statel (id :: got))))),
              wrap (ge, fn gv => state2 (destMonitor gv, got, numDest gs, 0)));

  fun state2 (mdp, got, potential, noreply) =
    if length got = n orelse
      length got + noreply = potential then []
    else sync (choose (wrap (unord (portReceive dp),
                                fn (x, id) => x :: (state2 (mdp, id::got, potential, noreply))))
                      wrap (unord (portReceive mdp),
                            fn (DEST FAILED id, _) =>
                                state2 (mdp, got, potential, if member id got then noreply
                                         else noreply + 1))
                                | _ => state2 (mdp, got, potential, noreply)))))
                                 in
      statel []
      end;

Fig. 2. Implementation of procReply

IV. EXAMPLE 2: PROCESSING REPLIES

Recall the implementation of RPC Multicast of section II: clients are probably not interested in receiving a
destination port and a unit gview event from their requests. Instead, they would prefer to use a library
function to process incoming replies. We will define the function

val procReply : int * ('b rpcreply dest_port * unit gview event) -> 'b list

procReply takes an integer n and the result of a request call and returns a list of reply values. This list
will have length no greater than n, but may be less than n if fewer than n servers reply. Its implementation
is in Figure 2.

A thread executing the procReply function is in one of two states, depending on whether or not the
unit gview describing the query destination set has been built. Initially, the function is in state1; it waits
for either a reply to be received on the reply group dest.port or for the unit gview build to complete.
After the gview is completed, the function enters state2 and monitors for replies or failed query group
dest.ports. The variable got is a list of query dest.port port.id's which have been acknowledged in
replies. The integer potential is the maximum number of replies, while noreply indicates the number
of failed dest.port's that never replied.

V. CONSISTENCY AND MEMBERSHIP

In sections II and IV we saw two examples of DML programs, and the latter motivated the need for explicit
failure information. An important issue in systems that support consistency-preserving programs is the way
they order failure information relative to other communication in the system. In DML, for instance:
1. When a src.port fails, its failure notification is ordered after all receive events for messages sent through the port.

2. If two failures occur simultaneously, their failure notifications are ordered identically at threads that receive them.

3. Any receive event of a port group will be ordered consistently with respect to a failure notification of any of the group's ports.

Properties two and three will be important in the next example.

VI. EXAMPLE 3: REPLICATED STATE

Although the DML syntax does not provide for distributed references, they can be simulated using port groups. In this example, we describe an implementation of distributed references through replicated state; the value of the reference is replicated to provide fault tolerance and quick dereferences. However, updates are slow.

Define the type 'a dref.query to be

datatype 'a dref_query = DEREF of ('a, unit) port_group 
 | UPDATE of 'a;

Copies of the replicated state communicate through 'a dref.group.

type 'a dref_group = ('a dref_query, 'a option) port_group;
fun mkDGrp () = mkGrp TOTAL;

Each distributed reference has an associated dref.group; threads dereference and update by sending messages through the appropriate group. For instance, the dereferencing function !! can be defined by

fun !! g =
  let
    val replyGr = mkGrp FIFO;
    in
      portTransmit (#1 (mkSrc (g, NONE)), DEREF replyGr);
      portReceive (#1 (mkDest (replyGr, ())))
    end;

Threads can install instances of the replicated state by calling the function install, defined in Figure 3. The implementation of install is notable for several reasons:

1. Dereference queries receive replies from every copy of the replicated state. A more intelligent (and lengthier) implementation would ensure that only one (local, if available) copy replies to each dereference.

2. dref.groups are totally ordered, so updates are received in the same order at each copy.

3. Updates and new dest.port creation events are ordered with respect to each other. This ensures that a new replica R processes the same stream of updates that old replicas process after observing R's creation.

4. There is no method for bootstrapping the example; in a complete implementation, there must be a function for creating the first replica.
fun install g =
  let
    val xg = mkGrp ();
    val (dest, view) = mkDest (g, SOME xg);
    val mdest = portMonitor view;
    fun server value =
      OrdEvt.sync (resolve (OrdEvt.wrap (portReceive dest,
        fn (DEREF g') =>
          (portTransmit (mkSrc (g', ()), value); server value)
        | (UPDATE x) => server x),
          OrdEvt.wrap (portReceive mdest,
            fn (DEST_CREATE (SOME g')) =>
              (portTransmit (mkSrc (g', ()), value); server value)
            | _ => server value))));
  in
    server (OrdEvt.sync (portReceive (#1 (mkDest (xg, ()))))
  end;

Fig. 3. Implementation of install

VII. CONCLUSIONS

We have illustrated that explicit failure information, coupled with the appropriate ordering properties, permits a flexible and convenient notation for high performance asynchronous communication.

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Implementing ML on the Fujitsu AP1000

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Abstract

The CAP ML project seeks to develop a version of ML that is suitable for use on a distributed memory multiprocessor architecture such as the Fujitsu AP1000. Language extensions are proposed that have been developed in conjunction with a programming methodology that is appropriate to that of a massively parallel computer whilst retaining a functional style. The implementation, which is based on the SML/NJ compiler and the SML2C compiler, is in progress. This paper focusses on the language design.

Introduction

When one contemplates implementing a functional language on a parallel computer, what first comes to mind is the well known advantage claimed for functional languages, that freedom from side-effects allows multiple processors to be utilised for the parallel execution of multiple arguments in any function call. There is also a long history of applicative language compilers which take advantage of the property of referential transparency, for a variety of purposes. For example, the implementation technique of graph reduction depends on it for achieving speedup through concurrency. However, experience indicates the opposite - it is hard to harness a multiprocessor for ML.

There are several lessons that have been learned that are important background as we embark on the enterprise of providing a functional programming capability on a machine such as the AP1000.

- Purely functional languages are not found suitable for large application programs. Lisp and ML are far more widely used than Miranda; although the latter has been widespread for some time, its niche in the market is in the education sector. Programmers still find important uses, in major applications, for global data structures that are updated, and for i/o operations that can't readily be characterised in functional terms.

- Although ML and Lisp both allow assignments to variables, the use of this feature is discouraged in situations where concurrency is expected. (ML discourages any such use by the very syntax of reference operations.) Thus the programmer should seek to structure a program so as to have as few routines as possible that side-effect the state.

- It is only sensible to spawn a process if it is likely to survive for a time which is long compared to its setup time. Experiments where a compiler does its best to recognise which sets of subexpressions can be executed concurrently show there is surprisingly little chance that substantial parallelism can be achieved in that way; some estimates suggest that typical Lisp programs would be unlikely to make effective use of just ten processors.
• The conclusion of all implementors of Lisp and ML compilers for parallel machines is that the
programmer should design algorithms with concurrent execution in mind and give explicit
advice about which parts of a program can be usefully mapped to processes. A common
design decision is for the system to spawn processes only at places where the user advises
that multiple arguments to a function should be evaluated concurrently.

• Most parallel Lisp and ML compilers have been implemented on shared memory multipro-
cessors and really depend on this fact for their success. The rule-of-thumb that has been
suggested is that each process should run for some thousands of instructions to have a cost-
effective existence. In many applications a programmer is likely to find many appropriate
situations.

• In a distributed memory machine, each processor has its own memory and so the setup cost
includes identifying (by following pointers) all relevant cells, copying these across a network,
and initialising a new heap space with this data. The operation is something like a garbage
collection so we would expect it to sometimes be most economical to simply copy all of the
heap space from one processor to another. In the distributed memory case, spawning processes
is likely to be appropriate only if such processes last for some millions of instructions, hardly
a function call.

Fujitsu AP1000 Cellular Array Processor

The AP1000 is a highly-parallel scalable computer with distributed memory. Each cell consists
of a SPARC CPU, a Weitek FPU, custom message controllers, and 16Mb of local memory. It
has a front-end host Sun4/330 which connects to the cells. These in turn are connected by three
separate high-speed networks - a 2D mesh-connection torus network, a broadcast network for one-
to-n communications, and a synchronisation network. Typical sizes are 64, 128, 256, 512 and 1024
cells.

This new style of expensive machine is of the same class as the Thinking Machine Corpora-
tion's CM-5, Intel's iPSC and Touchstone Delta. They share the common features of difficulty of
programming and paucity of software (beyond the C and Fortran compilers). The vanguard of
applications are those based in the numerical analysis of scientific problems. In these applications,
processors are typically allocated to independent calculations or computations on a hunk of an ar-ray. Typical techniques are Monte Carlo simulation and Finite Element methods, just as employed
on SIMD machines.

Although MIMD machines exemplified by the AP1000 apparently have more scope for fast
general purpose computation, we must learn how to use them less for Physics problems and more
for AI (mathematics, knowledge, reasoning etc.).

Language Design

Fine grain concurrency, especially of the sort envisaged in graph reduction, is inappropriate where
there is no shared memory. Because functions are closures, there will be potentially large amounts
of heap space that must be copied from one processor to another, even for quite short expression
evaluations. We see very coarse grain parallelism as necessary for the combination of distributed
memory machine and applicative language.
It is expected that the programmer will carefully design algorithms with concurrency in mind and will take complete charge of the processes, both as syntactic objects and dynamically executing entities. Since we insist that the extensions to ML should retain the applicative flavour of the standard language as much as possible, processes will take arguments and yield a result.

The AP1000 style of architecture imposes considerable overheads on process creation, and thus we adopt a programming methodology that discourages the use of more processes than there are processors. The programmer should aim to create the required processes early in the execution of a program and expect that they will last for a time that is long compared to the process startup time.

In order that long-running processes can be used successfully, we must allow them to cooperate by passing information. In the case of the AP1000 this can only mean we provide communication between processes by message passing or by distributed shared memory; currently, we have chosen message-passing as the most efficient system to implement.

This design of the language is intended to support a two-level style of program structure where the top level is the initiation of 'actors' that interact with each other by message passing. Within these top level processes, the programming should resemble that of whole ML programs, where I/O is replaced by message traffic among the 'actors'. Use of messages is certainly not referentially transparent but the careful programmer can still structure each process in the applicative style and write most component functions to be side-effect free.

Based on these major design decisions, we present paraML, an extension of Standard ML that we claim is suitable for programming the highly parallel computers of the future. The AP1000 is a leading example, being a machine with many powerful processors, each with its own large local memory. The major extensions to Standard ML are given below.

**Processes in ParaML**

We make changes to two areas of ML to accommodate our notions of processes. We add a new sort of declaration (ie process definition) and two new forms of expression - one to create a process instance and one which gets the value computed by the process.

**Process Definitions**

The ML code that is the abstracted form of a process is declared in a way that is very similar to a function definition. The difference is that the external view of a process must reflect the fact that the process can receive messages on named channels. Thus the type of a process will have the form similar to \( \alpha \rightarrow \beta \rightarrow \gamma \) where \( \alpha \) is the type of the argument supplied, \( \beta \) is the type (a record type) of the n-tuple of channels, and \( \gamma \) is the type of the result of executing the process to completion. The complete syntax is given in Bailey [1] and a forthcoming manual, but the following is the usual way in which a single process form is defined.

```
declaration:
    define pdef (channel ch1, ch2, ..., chn) pat = exp;

binding:
    val pdef = prd: 'a->'b ->> 'c
```

pdef is bound to the process form described in the definition; it is an ML routine that takes an argument matching pat, that produces a result by evaluating exp and that receives messages...
on the channels ch1, ch2, etc. pdef is called the process definition identifier. The words define and channel are new reserved words for ML and ->> is a new type constructor that is used in expressions for the types of process forms and process instances.

Of course, some processes that we wish to define will read no messages and so have no need of the channel list. However, this must be signified, like the unit argument to functions. The following syntax is appropriate in this case.

```
declaration:
define pdef nochannels pat = exp;
```

The indicator, nochannels, is a new reserved word. Finally, there may be several process definitions written in the same process form.

```
declaration:
define pdef (channel ch1, ch2, . . . chn) pat1 = exp1  
pdef (channel ch1, ch2, . . . chn) pat2 = exp2  
and pdef2 nochannels pat exp;
```

### Process Creation

Each instance of a process is created in a create expression; the code associated with a process form is applied to the arguments supplied (which must, of course, be of the right type) and a running process is then in existence until the expression in the selected clause yields a result. This newly created process executes concurrently with the process containing the create expression. Although we introduce the notions with a simple, but very typical instance, the full syntax can probably be inferred.

```
create p = pdef expl in exp2 end;
```

The syntax is intentionally close to that of local declarations, since the scope of the process identifier, p, is just exp2. pdef is a process definition identifier and expl is the argument that will be bound to the formal parameter of the process program. After this binding, process p is active and executes concurrently with the evaluation of exp2 (called the body of the create expression). The value that results from evaluating exp2 is deemed to be the value of this create expression.

Within its scope (exp2 in the above example), a process identifier is taken to be of type α ->> β, where α is the type of the channel record and β to be the type of the result.

It is possible to initiate multiple processes in the one create expression as the following example indicates:-

```
create p1 = pdef1 expl  
and p2 = pdef2 expl2  
in exp3 end;
```

### Getting Results

In a create expression, the process identifier's scope is the body (of the create expression) so that it can be used to reference the result of the process (when that result is available), to send messages to the process and as a process descriptor that can be passed to other processes.
There is a polymorphic operator called `result` which takes a process as its argument and yields the result that was produced by that process; `result(p)` will not return anything until process `p` has terminated.

**Message Passing**

The way to send a message to another process is by invoking the predefined function `send`:

```plaintext
send p #> c exp
```

In this example, `p` is a process identifier and `c` is one of the channel names of the process form of which `p` is a process instance. The construct `p #> c` has type `α channel` and for type consistency the message expression should have type `α`. The semantics of the expression is that the object that `exp` evaluates to is sent on channel `c` to process `p`. The type of send is `α channel → α → unit`. The sending process is not blocked. The only possible exception that can be generated by a send is where the process `p` has terminated. If process `p` does not have a channel `c`, then the error is detected by the type checker.

The messages that are sent to a process on one of its channels are extracted from that channel (a message queue) by the predefined function `get`, an example of which follows:

```plaintext
expression:
  get #<c
result:
  x: 'a
```

In this case, `c` is a channel of the current process, the type of which must have been `α channel`. Both `#>` and `#<` are new operators, chosen to resemble the IO redirection of UNIX.

**An Example**

The Sieve of Eratosthenes (SOE) is a classic problem with various solutions being algorithms that are capable of efficiently using a large number of processors. In the solution below, we have a pipeline of processes, each one of which takes care of the selection of one prime number. A sequence of all odd numbers which is fed into one end of the array, is filtered as it passes along, so that the sequence that goes to the `nth` process contains no multiple of any of the first `n` primes but contains all other members of the original sequence. When the sequence is reduced to nothing a message flows back the other way, gathering primes as it goes.

```plaintext
val sieve (channel data:int) ()
  = let prime = get #<data
      in if prime = -1 then nil
         else create s = sieve ()
          in let fun f -1 = send s#>data -1
              fun f dv = if dv mod prime <> -1
                 then (send s#>data dv; f(get #<data))
                 else f(get #<data)
              in f(get #<data);
```

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The main program is the following function:

```
fun soe(0) = nil |
soe(1) = nil |
soe(2) = [2] |
soe(n) = create s = sieve ()
    in let fun genlist g if g<= n
        then (send s#>data g; genlist(g+2))
        else send s#>data -1
        in genlist(3);
2::(result s)
end
end;
```

Methodology

There are a number of standard ways of structuring parallel programs, such as worker farms, space partition, data partition etc. We have coded examples of these to show that an application that is amenable to solution by one of these strategies, can readily be written in paraML without the programmer worrying about process creation and inter-process communication.

There is insufficient space available to properly discuss the various recipes, so the reader is asked to watch for a subsequent paper.

Implementation

The SML of New Jersey compiler was used as the starting point for the implementation of paraML on the AP1000. It is incomplete at this stage although sufficient of the task is done to have uncovered some interesting problems. These problems and their solutions are presented in [1] and will also be addressed in a forthcoming report.

References


Verification of Concurrent Systems in SML *

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Abstract

There can be different views of a concurrent, distributed system, depending on who observes it. The final user may just want to know how the system behaves in terms of its possible sequences of actions, while the designer wants to know which are the sequential components of a system or how it is distributed in space. Moreover, there is no widely accepted semantic model for concurrent systems.

In this paper we describe the use of the SML language in the implementation of a parametric verification tool for process description languages. It allows symbolic execution of processes at different levels, and provides facilities for equivalence checking.

1 Introduction

In this paper we describe the use of the SML language in the implementation of a parametric verification tool for process description languages (PDL). In the recent past, many verification tools for distributed concurrent system, which are based on process description languages, have been proposed (for a detailed survey see [6]). Any of these existing tools is based on a specific theory and, therefore, it is suitable for analysing a particular class of problems. In particular, all the existing tools are based on the so-called interleaving models.

The tool we have realized supports more than one concurrency model and allows the user to simply switch from one model to the other. Roughly speaking, it is a parametric tool that permits to observe many aspects of a distributed system. Among the others the temporal ordering of the events and their causal or spatial relation can be studied. An introduction to the general motivations and to the theory underlying our approach is given in [8].

The main motivation underlying our approach is that there is not only one kind of observer of a system. More precisely, there can be different views depending on the kind of observer. The final user of the system may just want to know how the system behaves in terms of its possible sequences of actions, in this respect the interleaving semantics is enough for him. He looks at the system as a black box.

On the other hand, programmers who have to implement systems usually think in a truly concurrent way: two concurrent processes are two concurrent processes not a single non-deterministic one. Also in debugging a big system this way of thinking is more convenient, one truly concurrent run of a system gives the same amount of information to a programmer than an exponential number of interleaving computations. Moreover, there is no widely accepted truly concurrent model, but there are many of them. A tool allowing parameterization with respect to various models would also help in understanding which model is more useful and would allow comparisons among them.

Parametrizing a tool in order to catch many different aspects of a distributed concurrent system requires:

- to have a common description language for the various aspects to be considered;
- to have a very fine description of the system at hand, from which all the necessary information for a large number of models can be extracted.

In order to satisfy the two items above we proceed in the following way: we first start from a very concrete representation of the system under specification in terms of a particular transition system (in the sequel denoted by Tcsc), whose transitions are labelled by their proofs. On this very detailed description

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it is then possible to define abstraction functions, called observation functions, which permit to recast the chosen observational model from the Tccs structure by throwing away some information on the internal structure of the studied system. As an example, we can define an interleaving observation function which, given a Tccs computation, i.e. a sequence of transitions, returns the computation as observed in the chosen model by forgetting, for any transitions, the whole proof but retaining the label of the performed action. Analogously, it is possible to define functions that permit to retrieve all the common models presented in the literature.

Once the interesting aspects to examine, i.e., once a specific observation has been chosen, one of the most widely used facility concerns the ability of proving behavioural equivalences among communicating systems. These equivalences, usually called bisimulations, assess that two systems are equivalent if, whenever the first may perform a (possibly complex) activity, the other one may as well, reaching states that are again bisimilar.

According to the purpose of having a very flexible tool for the analysis of distributed concurrent systems, the implementation is modular and open. By open we mean that an expert user can define his/her own observations and equivalences, and then use the standard facilities provided by the tool. We can see the tool as logically divided into two parts: a kernel and a library. The kernel contains the definitions of the input formalism, the functions to deal with simulation of processes and the functions to deal with the observed behaviour of the processes. Also the interface facilities are contained in the kernel. This part of the tool cannot be modified by the users and it defines also the interface that any observation module has to exhibit. More precisely, each component module of the library must contain the definition of an observation function and of an equality function between the observations of two computations.

To this respect the choice of SML as implementation language has been very convenient since SML allowed an easy design and manipulation of the various modules. Since parametricity and open-endness are two strong requirements for our environment, the modules system of SML turned out to be crucial. On one side, it forces the user to obey to some constraints in defining, from scratch, his/her own component through the notions of signature and structure. On the other side, the user can take advantage of the already defined components by defining a new component as composition of previously defined ones, through the notion of functor. Other important motivations to the use of SML have been the high portability of the language and the high number of existing tools which are implemented in this language. Indeed, a current work is the study of the possibility of the integration of our tool with other existing tools in order to reuse, when possible, already implemented software.

In the following, we describe the architecture of the tool trying to outline the parts which have most benefitted of the use of SML. Then we discuss our experience in using SML.

2 Underlying Theory

In this section, we will briefly recall the main definitions and results of the underlying theory the tool is based on.

An extended explanation of the underlying theory can be found in [3], where a general methodology for the definition of concurrent systems semantics is presented. This methodology is here instantiated with the language CCS [9].

In its general lines, the approach consists of four steps:

1. Define a transition system that captures the operational behaviour of the system. This operational description has to be very concrete, i.e. it has to capture all the information about transitions that the language is intended to describe. It is very natural to associate to CCS a concrete transition system, since the only information that a transition can carry on is its own proof. Hence, we choose as the basic level of description for CCS the proved transition system, which is the standard transition system of CCS, where the labels describe the proofs of the transitions.

Proved transition systems have been introduced in [1], and its algebraic versions in [10, 4]. The proved transition system for CCS is here called Tccs.

2. Build the computations of the system as paths in the transition system, and structure them as an observation tree (ordered them by prefix). The computations of the transition system Tccs are just sequences of (proofs of) transitions. Once ordered by prefix, the computations an agent can perform generate a (fixed) tree-like structure.
3. Define what are the observations of computations. These appear as labels on the nodes of the observation tree, describing the relevant aspects to be considered.

To observe a computation means to abstract away some details of the operational description that are hidden to the particular observer. For example, the interleaving observation is defined by abstracting away all the information in the proofs, recording only the actions of each transition and the order in which they are performed. In this way, to each computation is associated a sequence of actions.

Observational theories of concurrency have been presented in many places, the approach taken here mainly follows [2, 5, 4].

4. Define an equivalence between observation trees, based on the observations defined in step (3).

The equivalences that can be considered on observation trees are similar to those considered for transition systems. The so-called bisimulation equivalences [12] take into account the nondeterministic nature of processes, and hence are very natural for a theory of concurrent systems. Intuitively, two systems are bisimilar if, whenever the first may perform a (possibly complex) activity, the other one may as well, reaching states that are again bisimilar.

Different bisimulations can be found in the literature, here weak, strong and branching bisimulations are considered. Also trace equivalence is considered, which identifies two processes if they can perform the same sequences of actions.

In [3] a sound and complete axiomatization of many bisimulations for observation trees has been given. These axioms give the basis for a rewriting-strategy to the bisimulation problem, and they are used in this tool for implementing a rewriting decision procedure for strong bisimulation.

3 The System: architecture and implementation

In this section we describe the logical architecture of our tool. The architecture is illustrated in Figure 1.
Each box in Figure 1 represents a logical step that a generic user must perform to check the equivalence of two different distributed concurrent systems. Roughly speaking, these boxes are the kernel of our tool. Instead, dashed lines represent possible interactions with other entities like other tools or libraries.

The first step to be performed is to provide the description of the system to be manipulated in CCS. Then, the tool translates the CCS specification in the corresponding Tccs program. Tccs implements our basic syntax and it is the most concrete observation we provide to the user. Since we want to give to the user the possibility of defining his/her own observation algebras, we have equipped this module with a signature. Such a signature says what is exported outside the module, and therefore which are the offered functionalities. We register in the labels of the transitions all the information associated to it, i.e., their proofs in the SOS system defining the operational semantics of the language [13]. Some of the functionalities of this module are to check if a process has deadlock properties and to provide the string representation of processes. Up to now, we use CCS [9], but it is easy to adapt Tccs to support other formalisms by interacting with general compilers of specification formalisms.

The second step concerns the selection of the desired observation from a library. Also in this case, a signature which specifies the general characteristics of observation algebras is provided. The signature specifies which is the structure of agents, transitions and computations after the corresponding structures of the Tccs have been observed by means of abstraction mechanisms. The used abstraction mechanisms are the observation functions which take a Tccs computation and return the corresponding observed computation. All the modules which describe observation algebras must have this signature, therefore the functionalities which appear in this signature are parametric with respect to the chosen algebra. This interface contains all the functions which interpret the constructors of the datatypes agents and transitions. Moreover, the observation algebras must contain functions to perform the sequential composition and the equality of two computations. Note that this is not the syntactical equality function: for instance if the computations are partial orders, it implements an isomorphism checking. Moreover, a function to provide the string representation of observed transitions must be given.

Once the observation algebra has been fixed, the description of the system at hand is translated in a tree structure (the so-called observation tree) which permits to execute equivalence checking. The functionalities provided over observation trees are the equality of two trees up to associativity and commutativity, the building operator starting from an agent, an operator to reduce the size of these trees up to strong bisimulation, and finally a rewriting strategy to test strong equivalence of two trees.

Also a functor that lifts the basic functionalities of the tool to the level of observation algebras has been implemented. In other words, it is possible to look at transitions and computations of a system in the particular observation algebra selected. Analogously for the properties like deadlock and reachability. This is a very important module, since it permits to realize the parametricity with respect to the observation. Especially in this part has been crucial the choice of using SML. Indeed, the polymorphism of the language has permitted to simply recast the theory in ML thus obtaining the wanted results.

We are now ready to check equivalences. Our tool supports trace equivalence and three bisimulation-based equivalences (strong, weak and branching). All the equivalences are checked by an adaptation of the Paige-Tarajan algorithm based on partition refinement [11]. Finally, also a rewriting strategy to test strong equivalence has been implemented.

Some comments are in order for the library of observations. The user can specify which observation (s)he wants to use in three ways. The first possibility is by choosing an already existent observation from the library. The second one is by defining a new observation algebra from scratch as a new module in SML. This definition follows some standards: it has to fulfill some requirements established in a signature. Once this algebra is defined, the user can choose it from the menu and use the algorithms and facilities with this observation. While giving a definition, previously defined observation algebras and possible reuse of some modules (for example the structure LAB of labels) can be of help to the user. The last possibility is to apply an operator to the observation algebras that have already been defined. At present, the operation we allow over observation algebras is the product, which takes two observations and compose them in a new one. Obviously, the equivalence obtained in this way is finer than both the equivalences obtained with each observation. Indeed, it is finer or equal than the intersection of the equivalences. Many other operations could be defined over observation algebras, allowing the user to specify an observation domain as an expression in a language of observations. From the implementation point of view this simply means to define a functor which takes two structures which are observations and returns a new observation.

There is a wide spectrum of observations to choose from. The definition of new observations is also easy. In this way if a new theory is proposed within the methodology presented in [3], it is possible to obtain an automatic support for testing it with few efforts.
The tool is implemented in SML of New Jersey (version 0.66 of 15 September 1990) and runs over UNIX machines. Its code is organized into modules for ensuring a simple debugging and maintenance of the system.

4 Conclusions

Our goal was to implement a prototype of a parametric verification tool for concurrent systems based on a particular theory [7]. The main reason in choosing SML as the implementation language are the following:

- The compiler of NJ-SML is of free-distribution. This makes our tool portable, easy to distribute.
- NJ-SML is an high-level language adequate for rapid prototyping and its type discipline forces a nice style of programming.
- NJ-SML facilitates the interaction with many other existing tools already implemented in SML and permits to reuse code.

Since our main commitment was to stress the parametric aspects of the tool, NJ-SML has been particularly suitable to this purpose because of its algebraic nature. In a parametric setting to have signatures and correspondent structures allows us to establish the basic operation into a signature which can be instantiated with many different modules. Moreover, some of these modules can also be implemented by the user, once fixed the signature. The functors facility provides a very powerful way of implementing parametric modules with respect to an underlying algebra.

Besides these advantages, the fact of having non-functional constructs has been very useful since our system has a part which is algebraic in nature and also includes some imperative algorithms [11].

However, NJ-SML interpreter and compiler lacks of some important features. The version we have used is not well-documented and it does not provide facilities for developing interfaces, graphics and so on. It is also difficult to modify system parameters. Moreover, the error handling routine is poor. We think that all drawbacks are due to the absence of an integrated programming environment, which will be very helpful in prototyping in SML. A minor drawback is caused by the absence of overloading facilities which forced us to use too many different identifiers.

Acknowledgment: Giovanni Mandorino has implemented the algorithms for bisimulation testing and has participated in many discussions.

References

5 Appendix: A Very Simple Example

The input of the user is italic while plain text denotes the tool output. Comments are in *cursive style*.

The system displays a menu of observations. The user has to choose which observation(s) he wants to use.

**CHOOSE AN OBSERVATION ALGEBRA:**
- SPO $\Rightarrow$ Spatial Partial Orders Strong
- SPW $\Rightarrow$ Spatial Partial Orders Weak
- POR $\Rightarrow$ Partial Orders Strong
- POW $\Rightarrow$ Partial Orders Weak
- INT $\Rightarrow$ Interleaving Strong
- WEA $\Rightarrow$ Interleaving Weak
- STE $\Rightarrow$ Multisets Strong
- ALS $\Rightarrow$ Abstract Localities Strong

Once the observation algebra has been chosen, the system displays a menu with the bisimulation algorithms.

**CHOOSE THE BISIMULATION ALGORITHM:**
- S $\Rightarrow$ Strong Congruence
- W $\Rightarrow$ Weak Equivalence and Congruence
- B $\Rightarrow$ Branching Equivalence and Congruence
- T $\Rightarrow$ Trace Equivalence
- R $\Rightarrow$ Strong Rewriting

The system is now ready in a Weak Partial Orders setting. The user can now specify different agents.

\[
\text{val } p = \alpha \cdot \beta + \beta \cdot \alpha; \\
\text{val } q = \alpha | \beta; \\
\text{test}(p,q); \\
\quad \alpha \cdot \beta + \beta \cdot \alpha \text{ and } \alpha | \beta \text{ are not WEAKLY bisimilar} \\
\text{cho()}; \\
\quad \text{The user asks to change the observation.} \\
\]

\[ ...
\]

**WEA**

The Interleaving Weak algebra is chosen.

\[
\text{test}(p,q); \\
\quad \alpha \cdot \beta + \beta \cdot \alpha \text{ and } \alpha | \beta \text{ are WEAKLY bisimilar and WEAKLY congruent}
\]
A File System in Standard ML

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Abstract

This paper discusses the design and implementation of a distributed file system in Standard ML. The file system is implemented using persistent storage from the Carnegie Mellon Venari project, and communicates via Mach Remote Procedure Calls. This is work in progress, so details are subject to change, and no benchmarks are available yet. Performance figures will be available for the workshop.

1. Introduction

Many file systems have been implemented since the introduction of magnetic disk storage. To better organize files for humans, hierarchical directory structures are generally used. The implementation of these file systems has traditionally been done in relatively low level languages such as C. In Berkeley Unix,\(^1\) the user sees a single directory structure. Under the covers, however, Unix is dealing with disk blocks as untyped arrays of bytes. This is difficult and leads to code such as:

\[
\begin{align*}
\text{\#define itod(fs, x) } & \\
& (daddr_t)(cgimin(fs, itog(fs, x)) + \\
& (blkstofrags((fs), (((x) \% (fs)->fs->fp) / INOPB(fs)))))
\end{align*}
\]

After implementing such structures on disk, more code must be written to allow network access. In contrast, Standard ML permits a concise implementation of a very similar abstraction using ML datatypes, persistent storage, and Mach RPC to communicate. The ML implementation does not need to worry about superblocks, inodes, or cylinder groups — it merely sees one large tree to traverse. Since tree traversal is one of ML's strengths, the result is much more elegant than the above C example. The directory structure contains two mutually recursive types, File and Dirent. The root directory of a file system is initialized to DIRECTORY(ref [] : Dirent list ref), and so is a File.

```ml
datatype File = DATA of ByteArray.bytearray ref \\
| DIRECTORY of Dirent list ref \\
| LINK of string \\
and Dirent = FILE of string * int * time_t * File
```

The string in the Dirent is the file name, as expected. The integer is a unique ID assigned to the file at creation.

\(^1\)Unix is a registered trademark of Unix System Laboratories, Inc.

\(^2\)From /usr/include/sys/fs.h on a Mach 2.5 system.
2. Network Overview

In the 1990s, there is little point in designing a file system without taking network access into account. Since this is ML, a nearly stateless protocol is indicated. Inasmuch as there are only a few operations on a file, any stateless protocol will resemble Sun’s NFS [2] to some extent. For now, we adopt an NFS-like protocol, with the major difference being that our protocol communicates via Mach RPC instead of Sun RPC. With that said, the major RPCs and an informal definition of their semantics are as follows:

- **Lookup** takes a pathname and returns as a file handle the unique identifier for the file.
- **Read** takes a file handle, offset, and request size, and returns a string, along with the number of bytes read.
- **Write** takes a file handle, offset, request size, and data as a string, and returns the number of bytes it was able to write.
- **Mkdir** takes a pathname and creates an empty directory of that name if possible.
- **Rmdir** removes the empty directory named by its argument.
- **Create** creates a zero-length file named by its argument, if possible.
- **Unlink** removes its argument, if it does not name a directory, from the file system.
- **Makelink** makes its second argument a symbolic link to its first argument.
- **Readlink** reads the contents of the symbolic link whose pathname is passed to it.
- **Stat** returns the unique ID, time of creation, and file type of the pathname passed to it.
- **Readdir** takes a file handle, (which must point to a DIRECTORY file), and a magic cookie. It returns the directory entry corresponding to the cookie, and a new cookie, for the next directory entry. The cookie is implemented by using the unique IDs; the first file in a directory is read by passing in a special cookie. The same special cookie is returned to indicate the end of the directory.

Pathnames are passed as a list of pathname components; it is assumed that all but the last are directories, and the last component is a file or directory name. Since symbolic links are resolved by the client, they are stored as single strings. Since the actual definition of this interface is in MIG\(^3\), both C and SML clients can use it. A C interface, exporting Unix-like open, read, write, lseek, mkdir, rmdir, symlink, readlink, stat, and close functions would be an interesting exercise, especially for benchmarking purposes.

All calls handle symbolic links by returning an error code to the client, along with the number of the first pathname component that is a link. The client then does a readlink, to resolve the link, and retry the operation with the result. Since symbolic links should be resolved from the client’s viewpoint (in the case of multiple file systems), an interaction of this sort is needed. It would be fairly simple to optimize this interface by having the operation return the contents of the link along with the error code, thus avoiding an immediate readlink RPC. It is assumed that the client keeps a table of mounted file systems and their mount points; the client can find the longest pathname prefix to determine the correct file system. The client is responsible for lo'p detection. The client keeps a list of links it has traversed, and linearly searches this list for duplicates. While this approach may seem to have performance problems, based on my study of symbolic link usage on various Unix machines, links are relatively rare, and the vast majority of links point to their final targets. Thus, in most cases, link resolution takes constant time, while the worst case is \(O(n^2)\), where \(n\) is the number of intermediate links.

\(^3\)MIG is the Mach Interface Generator. It generates stubs for using Mach’s Remote Procedure Calls. It is roughly analogous to Sun’s \texttt{rpcgen}. While MIG originally only generated stubs for C and C++, Frederick Knabe and I reimplemented it in Standard ML to produce Standard ML.
After translating the above RPC definitions into MIG format, SMLMIG produces the following signature for the implementation of the file server:

signature NLFS =
  sig
    exception mlfs of int
    val lookup : MachIPC.port * string list -> int * int
    val read : MachIPC.port * int * int * int -> int * string * int
    val write : MachIPC.port * int * int * int * string -> int * int
    val mkdir : MachIPC.port * string list -> int
    val rmdir : MachIPC.port * string list -> int
    val create : MachIPC.port * string list -> int
    val unlink : MachIPC.port * string list -> int
    val stat : MachIPC.port * string list -> int * int * int * int
    val readlink : MachIPC.port * string list -> string * int
    val makelink : MachIPC.port * string list * string -> int
    val readdir : MachIPC.port-> int -> string * int * int
    end

Note that the last int return value is an error code, and the string lists are pathnames. The MachIPC.port is the file server's Mach port.

The client interface has two parts: one part executes the client half of the protocol, and the other is an application interface. The MIG-generated client interface is essentially the same as the server interface. The application interface wraps the RPCs in logic to handle symbolic links and errors, resulting in a Unix-like interface, but using exceptions to report errors instead of return codes and a global variable.

signature MLFSCLIENT =
  sig
    val pathSep : string;
    structure Common : COMMON
      type fh
    val fopen : string -> fh
    val read : fh * int * int -> int * string
    val write : fh * int * int * string -> int
    val mkdir : string -> unit
    val rmdir : string -> unit
    val create : string -> unit
    val unlink : string -> unit
    val stat : string -> int * int * int * int
    val readlink : string -> string * int
    val symlink : string * string -> unit
    val readdir : fh -> int -> string * int
    val reinit : unit -> unit
    val mount : string -> string -> unit
    val umount : string -> unit
    end

3. Implementation Details

The implementation of the unique ID scheme implies that a file system cannot have more than $2^{31}$ files. This is not considered to be a problem. A more natural unique ID would be nice, but there are no obvious candidates, since nothing is fixed in the file system. The file system is strictly a tree; Unix-style hard links do not exist. The absence of hard links is not considered a major drawback, since symbolic links are an effective replacement. While an inode-like structure could be used, it doesn’t seem to fit well into a high-level model.
of a file system. If the lack of inode-like structures becomes a problem, they would be relatively simple to implement and would not require a complete rewrite of the file system. The major changes that this would imply are the elimination of the separate unique IDs, and Dirent's would have an inode component instead of a File component. Presently, time.t is an integer, used to store the file creation date. Symbolic links are not interpreted by the server, so they are stored as simple strings, presumably parseable by the client.

The allocation system for the file system is one of its more interesting features. Files are allocated sequentially, until the heap fills up. At this point, the heap is garbage collected. Since the garbage collector is a copying collector, this corresponds to disk defragmentation in a traditional file system. A generational strategy for garbage collection will be very effective, as file lifetimes tend to be either very short or very long — consider, for example, temporary files made by compilers and text editors, vs operating system binaries and mailbox files. It may be necessary to periodically run the garbage collector, as a full scan of a large heap is bound to be slow. Along with the lack of inode-like structures, this allocation scheme is an experiment that may or may not succeed.

4. Low Level Details

The unique ID scheme uses a simple hash table that is updated by the create, mkdir, unlink, rmdir RPC's, and used by the lookup RPC. Since unique IDs are consecutive integers, the identity function is a perfect hash function, modulo hash table size. The hash table resolves collisions by chaining; this was very simple to implement in ML.

The persistence mechanism for the file system is based on the Venari project's work. In turn, this code is based on Mashburn and Satyanarayanan's Recoverable Virtual Memory (RVM) work [4]. The Venari persistence work exposes two separate heaps to the ML programmer: the traditional volatile heap, and the persistent heap[3]. Here the persistent heap contains the file system and little else, while the volatile heap serves SML/NJ's needs for runtime storage. A key performance issue for persistence is how much data is written out to disk. Since ByteArrays cannot be extended, writing past end-of-file requires that a new, larger ByteArray be created, data copied, and new data written. The current system does not allocate any space on file creation, but allocates 1024 bytes on the first write to a file. After that, the files grow exponentially when writing past end of file. Directories were made to contain references to the list of directory entries to minimize the amount of directory copying necessary.

Once the file system itself has reasonable performance, it is time to turn attention to network performance. In general, of course, large transfers traverse the network faster than small transfers, just as with disk transfers. The problem, however, comes when IP has to fragment packets. NFS, by default, sends and receives 8 KB blocks. Since Ethernet has a maximum packet size of 1500 bytes, each NFS transmission becomes 6 fragments. While this is acceptable on a single Ethernet, NFS does not work well across bridges or routers, where losing one fragment is fairly common. Reducing the transfer size to fit within 1 network packet provides a substantial performance increase in this case. While MTU (Maximum Transmission Unit) discovery would be the right thing to do, it is very difficult in general. Since IP requires networks to have a minimum MTU of 576 bytes, picking a transmission size of 1024 bytes seems fairly safe. In the common case of routing between Ethernet's, there will not be any fragmentation. In the worst case, there will be at most two fragments, so there should be a reasonable chance (certainly much better than with six fragments) of all the fragments getting through the bridge/router. Note that FDDI has a MTU of about 4500 bytes, and in general, increasing network speed seems to bring increasing MTUs, so while a RPC size smaller than 1500 bytes will become less optimal, the performance problems of packet fragmentation are always avoided. Since the page size of most modern workstations is in the 4-8 KB range, a 1 KB network transmission should give a nice readahead effect, assuming the file is being read sequentially from start to end, which is extremely common.
5. Future Work

One of the key benefits of this file system implementation is its brevity. The entire file system is less than 1000 lines of user written code, so it is ideal for performing file system experiments. Tuning the file system is the first priority:

- The best base for the exponential growth of the ByteArrays can be found experimentally. The tradeoff between byte copying cost and wasted space needs to be determined, and perhaps made adjustable on a per file system basis.

- An alternative scheme to support growable files would use a list of fixed length ByteArrays. While this could improve write speed significantly, read performance may suffer slightly due to the need to handle reads that straddle the block boundaries imposed by the server. Since reads tend to dominate writes, this is an interesting tradeoff.

- Pathname-to-handle lookups should be cached. This cache can have a hit rate of 70–80%.1

- The file server could be multi-threaded. If a slight decrease in reliability is tolerable, one thread could update persistent storage every 30 seconds, like the Unix update daemon. This could dramatically decrease the number of disk writes actually done by the file system for processes appending to the end of large files. The file server could also fork a thread (up to a limit) for each incoming request, in which case a multi-threaded implementation would change the Dirent type to add a lock per file. Depending on the desired semantics of multiple concurrent writers to a file, locks might only be used on directories. Locking would proceed with each thread holding two locks at a time, much like a monkey climbing hand-over-hand up a tree.

The major performance issue not addressed here is disk seeks. In general, controlling data placement on the disk is impossible from ML. However, it is worth noting that RVM uses a linear mapping from virtual memory address to disk block, so if it were possible to influence ML's memory use, then some control over disk layout would become possible.

After the file system is properly tuned, there are several interesting directions to take this research. One would be to follow the ideas of Gifford in [5], and make a content-addressable file system. With current RPC technology, search engines would have to be implemented entirely on the server side. The grep family and the Venari signature matcher could be integrated into the server without undue trouble, and one new RPC, mkdirbycontent, added.

The file system currently has no protection mechanisms. While Unix-style mode bits are useful to protect users from themselves, simple protection schemes do not provide any real security across networks. I believe it is better to explicitly offer no security rather than poor security – this way there are no nasty surprises, other than human error. The design of the file system does not preclude any access controls; again the conciseness of implementation favors experimentation with new authentication protocols.

Another interesting direction to explore would be to add transactions to the file system. Transactions could be both used to structure the file server, and made user-visible for users desiring consistent updates of multiple threads. The Venari project is currently investigating the semantics of transactions in ML.

6. ML Issues

The design and implementation of the file system raised two points about the ML language: there is little control over memory use, and there is no nice way to specify when non-pure functions (that is, functions that generate and/or observe side-effects) should be evaluated. For most application programs, memory usage isn't particularly important, but for systems programming it is vital. For example, if a cache is about to be
paged out to disk, it may make more sense to discard the cache instead. Consider a file system cache: there is no reason to write the cache out, as it is almost as fast to read the original blocks back in again. Standard ML does not offer this level of control, which would be quite useful inside an operating system.

The following fragment illustrates the function-application time problem:

```ml
functor rest(structure Netname : NETNAME
    structure RPC : RPC) =
  struct
    val file_server_port = Netname.lookup("localhost", "ML_File_Server")
    fun lookup filename = RPC.lookup(file_server_port, filename)
  end
structure T = Test(Netname)
```

Assume RPC is a structure containing client-side interfaces to remote procedure calls and Netname is the interface to a network name server, from which processes can check in or out interprocess communication ports. Clearly, T.file_server_port should be bound to the port the file server is listening on at runtime, not at functor application time, as presently happens. While the desired functionality can be simulated by judicious use of references, the result is not very elegant. With careful thought, it should be possible to provide this functionality without too much modification to the semantics of ML.

7. Summary

With Standard ML, SMLMIG, and the Venari persistence implementation, it is possible to implement a distributed file system. The implementation is concise and has reasonable performance. A variant of a common protocol is used for access to the file server, while the file server itself is implemented in a fairly unorthodox fashion.

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References

Implementing a Mixed Constructive Logic in Standard ML

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1 Introduction

In this paper we describe an implementation of a programming language MCL which is based both on constructive and classical logic. The language is intended for specifying, constructing, and executing verified programs. By combining elements of both constructive and classical logic, it is possible to eliminate some computational inefficiencies found in purely constructive type theories, but in a way that does not drastically change the style of proofs. The availability of both constructive and classical operations makes it possible to write specifications not available in other systems. We are working on a prototype implementation of the language in Standard ML [5].

2 Constructive type theory

In a Constructive Type Theory (CTT), proofs of constructive logic can be viewed as programs [2, 4, 7]. If F is a logical formula in a CTT, then a constructive proof \( \pi \) of \( F \) is a computation that stands for a justification, a piece of evidence sufficient for believing in the truth of \( F \). In the same way that an arithmetic expression stands for an element of datatype \texttt{int}, the proof \( \pi \) stands for an element of datatype \( F \). Thus in a CTT, proofs are programs, and formulas are datatypes.

Datatypes in Pascal, say, serve as very crude specifications of the programs that compute elements of the type. Formulas in CTT’s serve as more interesting specifications. For example, formula (1) below could serve as the specification of dividing \( z \) by \( y \).

\[
\forall x : \text{int}. \forall y : \text{int}. \exists q : \text{int}. \exists r : \text{int}. x = q \cdot y + r
\]

(1)

For the purposes of this paper we are not concerned about specifying more completely the relationship between \( z, y, q, \) and \( r \), although this is certainly possible. The two existential quantifiers here will suffice to demonstrate a source of computational inefficiency. In MCL syntax, formula (1) is written

\[
\text{all \( x: \text{Int}. \) all \( y: \text{Int}. \) some \( q: \text{Int}. \) some \( r: \text{Int}. \) \( x=q\cdot y+r \)}
\]

As a datatype, a universal formula acts like a function space, and an existential formula acts like a product. Hence an expression that has the datatype given by formula (1) is a curried function that takes two integers (call them \( x \) and \( y \)) and returns a nested tuple with two integer components \( q \)
and \( r \) and a final component of type \( x = q \ast y + r \) that specifies the relationship between \( x \), \( y \), \( q \), and \( r \).

Since proofs are programs, one can write translators for constructive logic. The basic idea of taking a proof of a formula and (say) interpreting it to produce a piece of evidence is the same as the basic idea of taking an expression and evaluating it to get a value.

Unfortunately, the most obvious way to implement a CTT causes runtime inefficiencies because uninteresting pieces of evidence are calculated. Consider the following proof/program of (1) written in MCL:

\[
\text{fun} \ x: \text{Int} \ . \ \text{fun} \ y: \text{Int} \ . \\
\langle x \ \text{Div} \ y, \langle x \ \text{Rem} \ y, e \rangle \text{ into some } r: \text{Int} \ . \ P \rangle \text{ into some } q: \text{Int} \ . \ Q
\]

In this proof we use \text{some-introduction} twice to exhibit the quotient and remainder. The expression \( e \) stands for the proof that the chosen quotient and remainder have the right properties. \( P \) and \( Q \) abbreviate the appropriate bodies of the existentials. The type-checker needs \( e \) to verify that the program meets the specification (1). At runtime, if the function above is evaluated for specific \( x \) and \( y \), then values for \( x \text{div} y \), \( x \text{rem} y \), and \( e \) will be calculated. If we are not interested in the actual piece of evidence denoted by \( e \) (and this is the usual case), then the evaluation of \( e \) is wasteful.

For our particular example, this is not much of a problem, but in the general case, evaluation of a function that justifies \( \forall x : t . \ \exists y : t' . F \) could spend an arbitrary amount of resource calculating a justification for \( F \).

The usual method [3, 6, 7, 9] for avoiding this problem involves modifying the specifications. Find the formulas whose proofs would cause undesired computations, then replace those formulas by formulas that are classically equivalent but have only computationally uninteresting proofs. (The proofs are uninteresting in the same sense that non-\( \bigcirc \) expressions of type unit in SML are uninteresting.) Modify the implementation so that it detects and optimizes away these uninteresting proofs. The result is more efficient execution.

Unfortunately, this technique has the side effect of making these more efficient proofs harder to read. For example, take the specification (1) above. If all we are interested in is \( q \) given \( x \) and \( y \), we could rewrite (1) as follows:

\[
\forall x : \text{int} . \forall y : \text{int} . \exists q : \text{int} . \neg \forall r : \text{int} . \neg (x = q \ast y + r)
\]

Proofs of (2) are classically equivalent to proofs of (1), but such proofs are also less clear.

3 Mixed constructive type theory

A more direct way to improve the efficiency of the interpreter is to allow the programmer to indicate what parts of the program are computationally significant. This is like Hoare logic in which the program segments, which indicate the computational part, are entirely distinct from the assertions, which are important for the purposes of verification. For example, the assertions can refer to variables that do not appear in the program and have no meaning at runtime.

In MCL we are exploring a language in which the user specifies the computational content by using both constructive \textit{and} classical quantifiers and operations in specifications. We keep \( \forall \), \( \exists \), and \( \forall \) with their constructive meaning, but now add \( \forall^* \), \( \exists^* \), and \( \mid \) as the classical counterparts. Now we can write the following specification:

\[
\forall x : \text{int} . \forall y : \text{int} . \exists q : \text{int} . \exists^* r : \text{int} . (x = q \ast y + r)
\]
which is computationally equivalent to (2) when uninteresting proofs of (2) are removed.

Programs in this logic can have two parts, a constructive part that actually denotes computations to be done, and a classical part that discusses but does not actually do computations. The rules of constructive logic are mimicked in the classical parts of the logic, but the classical parts of the proof also have access to rules of logic that are not constructively true. For example, to prove either $\exists r : \text{int} \cdot F$ or $\exists \bar{r} : \text{int} \cdot F$, you have to show how to calculate an appropriate $r$, but in the constructive version, this calculation must be computable; in the classical version, this calculation does not have to be computable.

MCL is based on the mixed constructive logic $S_1$ of [8] where a formal semantics can be found. The mix of classical and constructive formulas in $S_1$ makes it possible to write the formulas that are expressible when we mimic classical logic within constructive logic. Furthermore, it is also possible to write certain formulas that are otherwise inexpressible. For example, the formula $(\forall x : t \cdot \exists y : t' \cdot F) \Rightarrow (\exists y : t' \cdot \forall x : t \cdot F)$ is true in the semantics of [8], but is not known to be expressible in, for instance, NuPrl [2].

The rest of the paper is devoted to a description of MCL. We will concentrate mainly on syntactic and implementation issues here.

4 Language

In this section we give a brief overview of the language. The most important constituents of the language are the expressions/proofs and the types/formulas. We give a portion of the grammar for each of these two categories to suggest the basic appearance of the language.

Expressions consist of basic computational units like integer numerals, pairs and functions, as well as laws of logical deduction. Among the latter are the constructs for proving that something exists (some introduction), and the construct for reasoning about the existence of something (some elimination). All introduction and all elimination correspond to function abstraction and function application, respectively.

\[
\begin{align*}
\langle \text{expr} \rangle & ::= \langle \text{ident} \rangle \\
& ::= \langle \text{function symbol} \rangle \\
& ::= \langle \text{expr} \rangle, \langle \text{expr} \rangle \\
& ::= \langle \text{expr} \rangle.1 \\
& ::= \langle \text{expr} \rangle.2 \\
& ::= \text{injl} \ (\text{expr}) : \langle \text{type} \rangle \ \text{end} \\
& ::= \text{injr} \ (\text{expr}) : \langle \text{type} \rangle \ \text{end} \\
& ::= \text{case} \ (\text{expr}) \ \text{when} \ (\text{ident}) \ \text{then} \ (\text{expr}) \ \text{when} \ (\text{ident}) \ \text{then} \ (\text{expr}) \ \text{end} \\
& ::= \text{fun} \ (\text{ident}) : \langle \text{type} \rangle \cdot \ (\text{expr}) \\
& ::= \langle \text{expr} \rangle(\text{expr}) \\
& ::= \langle \text{expr} \rangle, \langle \text{expr} \rangle \ \text{into \ some} \ (\text{ident}) : \langle \text{type} \rangle \cdot \ (\text{type}) \\
& ::= \text{let} \ \langle\text{ident} \rangle, \langle\text{ident} \rangle : \langle \text{type} \rangle := \langle \text{expr} \rangle \ \text{in} \ (\text{expr}) \ \text{end}
\end{align*}
\]

These are not all the expressions in the language. In particular, certain acts of reasoning about integers, equality, and the like must also be represented by expressions in the language. For example, the expression $ax^2(e_1, e_2)$ is an element of a certain equality type.

The collection of types in MCL contains primitive types like Int, but most important, types corresponding to the logical connectives conjunction, disjunction, and implication. Conjunction
plays the role of the type of pairs. A more realistic programming language would contain more
general types for tuples and records, but they are not necessary for the purposes of studying the
efficiency of languages based on CTT's. Implication subsumes the role of the type of functions. It
is actually a special case of the "all" type. Quantifiers have both a classical and constructive form.

\[
\begin{align*}
(type) & ::= (ident) \\
& ::= (primitive type) \\
& ::= (expr) = (expr) \\
& ::= (type) \to (type) \\
& ::= (type) \mid (type) \\
& ::= (type) \lor (type) \\
& ::= (type) \& (type) \\
& ::= all (ident) : (type) . (type) \\
& ::= all* (ident) : (type) . (type) \\
& ::= some (ident) : (type) . (type) \\
& ::= some* (ident) : (type) . (type)
\end{align*}
\]

As in the language F-sub [1] there is a special type ?. The symbol ? can be used in place of a type
in hopes that the MCL system will be able to determine the correct type.

Here is the remainder example in MCL.

(fun x:Int. (fun y:Int.
  <(Div <x,y>), <(Rem <x,y>), (ax2 x,y)> into (some r:Int.(x=(Add <(Mul <q,y>),r>)))>
  into (some q:Int.(some r:Int. (some r:Int.(x=(Add <(Mul <q,y>),r>))))) ))

Here is the same example again. This time the MCL system is able to fill in some of the type
information.

(fun x:Int. (fun y:Int.
  <(Div <x,y>), <(Rem <x,y>), (ax2 x,y)> into (some r:Int.?))>
  into (some q:Int.??))

5 Implementation

In this section we describe the SML implementation of the MCL language. This basic approach
could be taken to implement any interactive language system and is instructive for that reason.
We show the top-level structure of the MCL system and show how SML can be used to make a
stand-alone system.

The function TopLevel does the so-called read/eval/print loop. In fact, it does six things.

1. Prints a prompt to cue the reader for input and reads a line of input.
2. Parses the input line. This is the function of parse. This creates an abstract syntax tree ast.
3. Analyzes the expression. In particular, checks the types of all the subexpressions. The function type_check performs this part.

4. Evaluates the expression to normal form.

5. Prints the expression, its type, and its normal form.


Here is a (slightly simplified) version of the top-level function.

```ml
fun TopLevel () = (  
  let  
    val input = read ();  
  in  
    let  
      val ast = parse (input);  
      val (e', tau) = type_check (InitialTypeEnv, ast);  
      val e'' = eval (InitialEnv, e');  
    in  
      output (std_out, "expr: " - (print e') - "\n");  
      output (std_out, "type: " - (print-type tau) - "\n");  
      output (std_out, "eval: " - (print e'')) - "\n");  
      flush_out std_out  
    end  
    handle e =>  
      output (std_out, "Exception raised: ", (System.exn_name e)) - "\n");  
  end;  
TopLevel ()  

(* Loop *)
```

Notice that exceptions during read, in particular interrupts, are not captured to allow the function TopLevel to be exited.

The parser was implemented using ML-Lex and ML-Yacc, a lexical analyzer generator and parser generator for Standard ML written by Andrew Appel, James Mattson and David Tarditi of Princeton University. This is a particularly convenient way to build elements of the appropriate datatypes. In this way full advantage can be taken of the ML language in representing and manipulating syntactic terms of a language without sacrificing some user-friendly concrete syntax. Although a yacc parser is not as simple as some LISP read macros, it is flexible, general, and based on well-understood concepts.

It is possible in Standard ML of NJ to create a stand-alone interpreter for a language by using the command exportML. The following code causes an executable image of the ML system to be written to the file mcl.

```ML
if exportML "mcl"
  then (  
    output (std_out, "MCL version 0.4\n");  
    output (std_out, " Type "C to leave MCL and get back to SML/NJ\n");  
    TopLevel ()  
  ) else ();
```

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The function `exportML` returns false when the image is created. The executable image is a snapshot of the system as of the time the function was called. Thus the image can be preloaded with all the functions required for MCL by defining them before the execution of the `exportML`. Executing the file `mcl` causes the ML system to be reactivated at the point where the image was created. However, `exportML` returns the boolean value true in the reactivated system. Hence, using the code above starts the top-level read/eval/print loop as soon as the saved image is executed.

References


Experiences with ML for Building an AI Planning Toolbox

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Abstract
The qwertz toolbox is a system of ML modules for implementing AI planners. We discuss why ML was chosen to implement the system, provide a few guidelines for using the language, based on our experience, and make some suggestions for language extensions and programming environments.

1 Background
The topic of our research is AI planning [1], i.e., constructing a course of action for an agent, given information about the current state of affairs and the agent's goals. Although planning work in general, and of course our own results (e.g., [3, 7]) are interesting, they will not be of further concern to us here. Rather, we focus on the reasons for our decision to implement qwertz, a planning software toolbox, in Standard ML, and make a few comments about the language based on our three years of practical experience with it.

The reason behind our choice of ML has to do with the state of the art in planning at the project start time in 1988. In those days, many researchers, including ourselves, felt the field suffered from a serious problem: there was an extensive literature about algorithms, systems, and techniques—such as nonlinearity, abstraction, or hierarchical planning—but this literature was typically insufficiently precise and complete to enable others to easily reimplement what was described. To quote T. Dean from a paper written then [6]:

Today, these basic and intuitively appealing methods (for classical AI planning) are known largely by association with archaic and poorly understood programs; these methods cry out for careful reexamination and precise formulation.

One of our project goals was to rectify this situation. We aimed to "rationally reconstruct" classical planners such as to facilitate their reimplementations by others, in their favorite language. Thus, clear design and readable code were of prime importance, rather than tricky efficiency. Also, we dreamed of finding a clear conceptual organization of planning methods which could be mirrored by an analogous modularization of our planning software: orthogonal methods should be implemented orthogonal modules (unless the methods only appear orthogonal).

So, setting aside the particular scientific problems of AI planning, we faced two general software engineering issues. First, architecture: How to build a toolbox which facilitates the combination of
orthogonal methods, and experimentation with alternative implementations of each method? And second, didactics: How to avoid the trap of understanding these methods ourselves, but failing to deliver a description of them sufficient to allow others to easily reproduce our results? To cope with these problems, we decided to use three techniques:

**Functional Programming.** We think that, owing to referential transparency, functional programs are usually easier to understand than imperative programs.

**Algebraic Specifications.** This approach composes well with functional programming. Moreover, it is conceptually close to the kind of mathematical models used increasingly within AI.

**Literate Programming.** Knuth's idea of programs as literature [4], intended primarily for humans to read, rather than for computers to execute, fits well with our purposes.¹

Our interest in functional programming and algebraic specifications lead naturally to our contemplating using ML. However, we were reluctant at first to take this step. Within AI generally, and also within our institute, Common Lisp has established itself as the dominant language. Presumably, the intended audience of our publications would be familiar with Lisp, but not necessarily ML. Nonetheless, we decided that the benefits outweighed the risks in our case, and chose ML.

The toolbox now contains a sizeable library of modules. Some may be of general interest, such as those for sets and streams, and often have multiple implementations. Example AI specific modules are those for symbolic expressions, theorem proving, reason maintenance and heuristic search. Of course, there are also functors for constructing AI planners. An experimental planner we have built "beautifies" business graphics, and communicates with a graphics editor written in Common Lisp. The qwertz toolbox will be made freely available sometime soon.

The rest of this paper explains the reasons for our preferring ML at the time, discusses some of the problems we faced, including organizational ones, and reflects about the wisdom of our choice after three years of actual ML use. We also take this opportunity to provide some feedback to those responsible for the evolution of the language, and those developing ML programming environments.

## 2 Why ML?

Of all the features which persuaded us to choose ML, three stand out: compile time type checking, MacQueen's module system, and ML's small size. The principal arguments against static typing have been: 1) type declarations are awkward in an interactive language; and 2) existing type disciplines prohibit an unacceptably large class of correct programs, making it very unwieldy to implement certain abstract data types, such as lists and other "containers". Type inference and polymorphism have solved both of these problems quite well. Some Lisp programmers familiar with ML still maintain that ML's type system is too restrictive, as it does not support union types. However, we have never missed this ability in practice. Although there is at least one Common Lisp implementation which supports static type checking, ironically this requires a large number of type declarations. We prefer ML's approach.

Perhaps it is not obvious why we should be so interested in compile time type checking, if our main interest is the clear description of a software system. Does the fact that a Lisp system checks types at runtime make a correctly typed Lisp program any less comprehensible to the human reader? Presumably not; but being able to catch all type errors before publication is very comforting to human

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¹We have developed an SGML-based system for writing literate programs in any programming language [2]. The system is available for free by anonymous ftp. Contact the authors for more information.
Regarding ML's module system, its main attraction is its obvious relationship to algebra. There is nothing comparable in Common Lisp. The language constructs which come closest to ML's signatures, structures and functors are Lisp packages and CLOS, the Common Lisp Object System. However, the differences are more numerous than the commonalities. Suppose one tries to compare structures to packages. There is a single, global naming environment for Lisp packages, which makes them quite inadequate for programming in the large. Suppose two programming teams inadvertently use the same package names? Without access to source code, there is no way for package users to resolve such conflicts. Even with source code, it is a thankless, tedious job. ML structure names are dereferenced at compile time, making it easy, usually, to find a loading order which avoids such problems. There are several other significant differences between structures and packages, such as the fact that structures may be organized hierarchically, but there is no space to discuss these further here.

A comparison of CLOS and ML's module system may be an interesting subject, but we can only scratch the surface here. The specification of the CLOS portion of Common Lisp alone dwarfs all of ML. For us, this is a major limitation. It would be too much to expect our poor readers to learn all of Common Lisp, including CLOS, before they can comprehend our system. Alternatively, we could have taken the trouble to define some subset suitable for our didactic purposes, but it is more convenient to be able to refer to a number of excellent, existing texts on ML, such as [5].

However, there are deeper reasons for preferring ML's module system. The object-oriented paradigm, at least as it is usually realized, unduly mixes specification and implementation levels. A class is not (only) a specification of an object, but also a generator of its instances. It serves the role of both signatures and functors in ML. Also, whereas membership of an object in a class is determined by a simple search of an inheritance graph, whether or not an ML structure satisfies a signature is determined by pattern matching, which is considerably more powerful and flexible. Finally, classes have elements of both modules and abstract data types, but are less than optimal in either role. This said, one advantage of CLOS, and other object-oriented systems, is the ease with which some abstract data types can be extended. We have a bit more to say about this below, in our wish list for ML.

There are a variety of other reasons why we preferred ML for our project, but there is only room to mention one more: ML is small, but not too small. It includes all the features we feel we need, such as exception handling and a powerful module system. Some complain about the minimal size of ML's standard library. But, for our purposes, this was more of a feature than a bug; it reduces the amount our readers must know before they can begin to understand our system.

3 Programming Guidelines

It is possible to write incomprehensible programs in any language, and ML is no exception. We have adopted a few conventions which we think make our toolbox easier to understand and use. These are only guidelines. There have been occasions to deviate from them. Only the most significant guidelines will be mentioned here.

Avoid Imperative Features. Presumably there is no need to justify this to a functional programming audience. But we are happy that ML does have imperative features. When published algorithms we needed are clearly imperative, we implemented them in the same way, rather than trying to invent functional alternatives.

Avoid Milner Polymorphism. This guideline may appear somewhat strange. However, modules provide a better way, in our opinion, to realize polymorphic types in most cases. Equality types and
the difficult system of weak type variables are two indications of problems with Milner's approach. Why should there be a special language construct for one abstract data type? We use an EQ signature instead, with an element type and an eq function, which may be any equivalence relation. To date, we have violated this guideline only once, in our STREAM signature for "lazy" sequences. How could flatten : 'a stream stream -> 'a stream be realised without Milner style polymorphism?

Do Not Always Use Closed Functors. Some consider it good practice to use only closed functors, i.e. to pass all substructures as functor parameters and to avoid free references to other functors. We cannot agree. The design of a functor should respect the needs of its intended users. In a functor for symbol tables, for example, another functor for hash tables may be used. Most users of symbol tables will be happy to delegate responsibility for choosing a suitable hash table implementation.

Declare Only Meaningful Types in Signatures. Types are "meaningful" only if some set of operations have been defined for them. In a signature for tables, e.g., it makes little sense to include a key type when this type is completely unconstrained. Instead, include some comparison operation for keys or include a Key substructure constrained by some appropriate signature.

Do Curry Function Definitions. This is another area where ML offers too much flexibility. If some functions are curried and others not, willy nilly, the poor programmer tends to forget which choice has been made, and the syntax of function calls tends to be an ugly mess. We chose, somewhat arbitrarily, to curry function definitions.

4 Surviving in a Lisp World

In the AI division of our research institute, the de facto standard language is Common Lisp. Our choice of ML has not made it easier to cooperate with our colleagues. Our original justification was that we were using ML to specify our system, implying we would someday implement the system in Common Lisp. We have been doing our best to avoid the inevitable. If you find yourself in a comparable situation, here is a tip.

To facilitate communication between Lisp and ML processes, in either direction, we have implemented an s-expression module for ML. This allows us to read and write most Lisp data structures. The concrete syntax of our planning language, for example, uses s-expressions in the traditional way. This makes it easy and familiar for others to use our planners as if they were working with a special purpose AI language embedded in Lisp. Moreover, we have written a simple CLOS class for creating and using instances of our planners. Lisp users need not know that the constructor function for the planner starts an ML subprocess. What they don't know won't hurt them.

The overhead incurred by sending s-expressions between Unix processes is negligible, at least for our application. Also, if the ML implementation has an application generator which removes all unreachable code, such as the compiler, this arrangement does not place excessive additional demands on memory. The runtime data structures we generate in ML would also have had to be generated in Lisp.

This approach is something of a crutch, to make up for the lack of some way to share functions and other values, regardless of which language they are written in. We have a few more words to say about this subject below.

5 Wish List

Let us take this opportunity to make a few suggestions to ML implementors. Some of them may not be very constructive, for which we apologize, as it is probably too late to do anything about them. The
comments are grouped into those concerning the language itself, and suggestions for future programming environments.

First the less constructive comments: We are not very happy with some aspects of ML's syntax. It is too complex, both to read and, especially, to write. The main sources of the problem seem to be the redundancy of curried and uncurried functions, as mentioned previously, and unfortunate interactions between infix operators and the left to right evaluation strategy. It is not always clear where one needs parentheses. Honestly, how many of you were not irritated that first time your ML compiler complained about a Boolean expression like foo x andalso bar x? If the left conjunct does not require parentheses, why should the right one?

This syntax problem brings us to another gripe. ML was originally developed because of some weaknesses of Lisp. However, incomprehensibly, several good features of Lisp did not find their way into ML, such as 1) built-in data types for symbols and symbolic expressions, for easy input and output of most structured data and, although we haven't needed this feature ourselves 2) the representation of programs as symbolic expressions, making it easier to manipulate programs as data. We have added symbols and s-expressions to our toolbox, but they would be easier to use if there were syntactic support for them, just as there are special syntactic means to construct numbers and strings.

Here is a constructive suggestion: The only feature of object-oriented programming which we sometimes miss in ML is the ability to extend data types across module boundaries. There may be numerous theoretical and practical difficulties which we do not appreciate, but perhaps it would be possible to modify ML's datatype construct as shown in the following example:

```ml
functor MyColors (C : sig
    datatype color = red | blue
    val paint : color -> unit
end) =

struct
    datatype color = C.color | yellow
    extend C.paint yellow = ...
end

Structures generated by this functor would have this principal signature:

```sig
    datatype color = red | blue | yellow
    val paint : color -> unit
``` end

As datatype declarations are already permitted in signatures, this suggestion does not appear to violate any abstraction barriers. The new key word, extend, is used instead of fun to continue the definition of a function to handle the new cases.

In object-oriented languages, an object is an instance not only of its parent, but also of all ancestors of its parent. Analogously, it would be useful if all structures matching the extended signature also matched the "parent" signature. To achieve this effect, the interpretation of data types would have to be modified to mean that the type includes at least, rather than exactly, the constructors listed. Failure to handle some constructor not mentioned in the signature, but part of the actual structure used, should presumably raise a runtime Match error.

Here is our final language suggestion. Using include it is possible to extend signatures. A comparable exclude declaration, for subtractive "inheritance", would be useful for minimizing dependencies in functor definitions.
Let us now turn to environmental issues. Our comments here are based on our experiences with SML of New Jersey, which we can highly recommend. But nothing is perfect.

First, SML of NJ requires enormous amounts of RAM. We fail to understand why ML should be any more memory hungry than, say, Macintosh Common Lisp, which is quite useable with as little as 2 MB of RAM. One reason for this problem may be that all modules used in a program need to be loaded at once, during development. When an application is "exported", memory demands can be radically reduced, due to dead code analysis and, of course, because the compiler is no longer necessary. The compiler may also be used in batch mode, but in this case it produces binary files in another, incompatible format, and does not perform type-checking across module boundaries. Our first wish would be an improved batch compiler without these shortcomings. To test programs, the required binaries could be loaded into a minimal environment allowing expressions to be evaluated, but no new functions or modules to be compiled. An interpreter should suffice. It would also be nice if there were some way to explicitly unload modules, to avoid having to restart ML when memory starts getting tight.

Finally, it would be helpful if an ML compiler would produce object code compatible with others languages. Ideally, the object code format would be the same as that produced by Unix C compilers. This may be the single most important improvement, if one is interested in increasing ML's popularity. Imagine gaining access to POSIX or X Window libraries just by defining appropriate ML signatures. However, if this is not feasible, it would be quite useful if at least the functional programming community, including Lisp and Scheme, would adopt some standard object code format suitable for this class of languages. Poplog, which supports mixed ML, Common Lisp, Prolog and Pop-11 programming, demonstrates this can be done, but it is not a standard.

6 Would We Do It Again?

Yes, definitely. Despite our complaints, we still are convinced that we made the best choice. No better alternative has appeared within the last three years. Nor are there signs of anything better on the horizon.

References


Attribute Grammars in ML

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1 Introduction

Attribute grammars provide a formalism for assigning meaning to parse trees of a context-free language [Knu68]. Because of their syntax-directed form and declarative style, they provide a useful notation for specifying compilers [KEZ82] and language-based editors [RT88]. This paper reports on a system we are developing, called AML (for Attribution in ML), which is an attribute grammar toolkit for building language-based editors using SML [MTH90].

AML is a spiritual heir to the Synthesizer Generator project at Cornell University [RT88], which focused on efficient incremental evaluation techniques, and the Pegasus project at AT&T Bell Laboratories [RG86], which focused on providing a high-level foundation for interactive systems. In our system we plan to build on the evaluation technology of the Synthesizer Generator while using a higher-level foundation for the implementation.

In the next section, we give some background about attribute grammars. Then we describe our specification language for attribute grammars, followed by a description of the internals of our system. Lastly, we discuss the project's status and future plans.

2 Attribute grammars

An attribute grammar is a context-free language (CFL) together with a set of attributes for each nonterminal and a set of attribute evaluation rules for each production. An attribute is either synthesized or inherited; for each production p : X₀ ::= X₁ ..., Xₙ, there are evaluation rules that define the synthesized attributes of X₀ and the inherited attributes of X₁, ..., Xₙ.

2.1 Attribute evaluation

Each node of a parse tree in the underlying CFL is labeled with instances of the attributes associated with the nonterminal at the node. The evaluation rules define a set of constraints on the attribute instances, and computing the attribute values can be viewed as a constraint solving problem. An attributed tree is said to be consistent if its attribute values satisfy the constraints defined by the attribution rules. The simplest way to solve this problem is to topologically sort the attribute instances by their dependencies and evaluate in topological order. For many grammars, however, more efficient and specialized evaluation strategies can be used. Attribute grammars are classified by their evaluation strategies; for example, the parser generator yacc implements a grammar in which all attributes are synthesized and can be evaluated in a single bottom-up pass. One important class of attribute grammars is the class of Ordered Attribute Grammars (OAGs) [Kas80]. OAGs

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are interesting because they include many useful "real-world" grammars, and because they can be efficiently evaluated by fixed-plan evaluators.

2.2 Subtree replacement and incremental evaluation

In his seminal thesis, Reps showed that attribute grammars provide a useful formalism for defining language-based editors [Rep82]. Such systems use attributed abstract syntax trees to represent programs being edited and map editing operations to subtree replacements (e.g., a delete operation is implemented by replacement with a null tree). After a subtree replacement operation, the tree will no longer be consistent; Reps and others have described so-called "optimal" evaluation algorithms for restoring attribute consistency [Rep82, Yeh83, Hoo87, Hud91]. Applications of this technology include structured editors for programming languages, interactive theorem proving systems for structured editors for programming languages, interactive theorem provers [Gri87] and code generators [Mug88].

3 Attribution in ML

Our approach to supporting attribute grammars in ML is a fairly conventional generator-based approach. Our system takes an attribute grammar specification and generates a tree machine for building and manipulating abstract syntax trees and an evaluator for the grammar. These are combined with grammar-independent modules to construct a complete system. We chose this approach over embedding the system in the interactive ML system, because it allows more flexibility for analysis and optimization. The remainder of this section describes our specification language.

3.1 The specification language

The AML specification language has two parts: a core language for specifying grammars and a module language for structuring large specifications into separate units. The core language is sufficient to specify an attribute grammar as a monolithic specification, but it is often useful to structure large specifications in a modular fashion.

An attribute grammar specification in AML consists of three main parts: the abstract syntax, the associated attributes, and the attribution rules. Chapter 4 of [RT88] gives a sample specification of a language-based editor for a simple imperative programming language; we use this example to illustrate our specification language.

The abstract syntax is declared using a limited form of the ML datatype declaration, which we call a prodtype declaration. For example, the abstract syntax of expressions can be declared as follows:

```plaintext
prodtype exp
  = EmptyExp
  | IntConst of int
  | True
  | False
  | Id of string
  | Equal of (exp * exp)
  | NotEqual of (exp * exp)
  | Add of (exp * exp)
```

This defines a nonterminal type `exp` with a set of productions. Prodtype declarations are more restrictive than datatype declarations in that the argument types of the productions (constructors) are required to be tuples of ground and nonterminal types.

The attributes associated with the nonterminals are declared separately from the abstract syntax. For example, the following declaration states that expressions have a synthesized type attribute and an inherited typing environment:
The last part of the specification are the rules for attribution. These require notation for naming
the nonterminals of a production and the attributes of a nonterminal. We use ML-style pattern
matching, with minor extensions, to bind names to the nonterminals and use the notation $X.a$ to
refer to attribute $a$ of $X$. The following are the typechecking rules for expressions in the simple
editor:

```ml
attr ex with
  syn typ : type_t
  inher env : (string * type_t) list
end
```

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editor:

```ml
attr eO : exp
  = EmptyExp
  | (IntConst _) with rule eO.typ = EmptyTy end
  | (True | False) with rule eO.typ = IntTy end
  | (Id id) with rule eO.typ = IntegerTy end
  | (Equal(_, _) | NotEqual(_, _)) with rule eO.typ = BoolTy end
  | (Add _) with rule eO.typ = IntTy end
  | (Equal(e1, e2) | NotEqual(e1, e2) | Add(e1, e2)) with
    val error = if (incompatibleTypes(e1.typ, e2.typ))
      then NONE else SOME "TYPE ERROR"
    rule e1.env = e2.env
    rule e2.env = eO.env
  end
```

In these rules, $eO$ is bound to the left hand side of the productions. Each clause defines a production
pattern with a collection of rules, which are ML expressions. We use the or-pattern notation to
group several productions that have identical rules (e.g., the last clause). The rules for a given
production can also be factored across several clauses (e.g., the rules for Equal). The val binding in
the last clause defines the local attribute error for those productions. Local attributes are associated
with productions, and are typically used to factor common subexpressions and, as in this case, to
communicate predicates about the node's state to external observers. For example, a pretty printer
might highlight a node's subtree when the error attribute is not NONE.

### 3.2 Modular specifications

Attribute grammar specifications tend to be considerable in size and non-trivial in complexity.
Attribute grammars as such do not provide any means for modularization. As pointed out in [Kas91],
an effective strategy is to allow a module for an AG to contain the attribution of one semantic
aspect only. Several approaches have been advocated to modularize AGs [DC88, FMY92, Far92]. While
these approaches address the problems of partitioning a grammar specification, they do not provide
the concept of a module.

Our approach is based on the module facility in ML. Grammar structures, akin to ML structures,
allow encapsulation and grammar signatures, akin to ML signatures, provide information hiding (i.e.,
the interface to the outside world). A grammar structure specifies a particular semantic aspect of the
problem being expressed as an attribute grammar. A grammar structure can have several grammar
signatures providing different views of the module. A relevant grammar signature is used by another
module to import exactly the information it requires. Grammar signatures allow information about
productions of the grammar and the attributes of the grammar symbols to be imported and exported
in a controlled manner, aiding comprehensibility and maintainability.

### 3.3 Computation with attributed terms

Since prodtype declarations in AML are just a restricted form of datatype declarations in ML, it is
straightforward to express computations on the unattributed view of a tree. This is especially useful
for supporting features such as syntactic reference (i.e., using a subtrc. as an attribute value) and higher-order attribute grammars [VSK89, TC90].

It is also desirable to allow user computations on attributed terms. To support this, we generate an abstract interface to the tree machine that provides for safe manipulation of attributed terms. Operations include attributing unattributed terms, reading attribute values and subtree editing (with retribution). This framework allows for a general and uniform platform for building applications that can manipulate attributed terms and allow access to attribute values.

4 Implementing attribution in ML

A number of issues must be addressed in the implementation of the tree editor and evaluator:

- Navigation and subtree replacement operations for attributed trees must be supported.

- It should be possible to convert between attributed and non-attributed versions of terms. Attribution of a non-attributed term is done by the evaluator; the converse should also be possible.

- A mechanism for associating attributes with tree nodes must be provided. It is especially useful for this mechanism to support sparse attribution (e.g., because of copy rules). In addition, operations for accessing and setting the values of attributes must be provided.

The Synthesiser Generator uses a heavy-weight tree-node representation that relies on mutable fields in the tree nodes (attribute instances, parent and child pointers, and other status fields). This representation supports efficient navigation, tree editing and attribute evaluation, but does not support sparse attribution, sharing of trees and easy mapping between values computed by user code and abstract syntax trees. Furthermore, the heavy reliance on mutable fields does not map well to ML, since it requires ref cells, which add extra space and garbage collection overhead.

Our approach is light-weight: we use the datatype equivalent of the prodtype declarations as our tree representation, and use paths to label nodes and to support navigation. Mutable state, such as attribute values, is held in auxiliary data structures that map tree nodes to their state. In this scheme, an attributed tree is just a pair of an unattributed tree and a map from nodes in the tree to their attribute instances.

4.1 Abstract syntax trees and paths

The prodtype declarations in an AML specification are translated to the equivalent ML datatype declarations. Because it is often necessary to refer to an arbitrary tree node, we generate type tree_t, which is the discriminated union of the nonterminal and terminal types. We also generate functions for examining, visiting and replacing the children of a tree node. These types and functions are contained in a structure that matches the following grammar-independent signature:

```
signature TREE =
  sig
    type tree_t
    exception Child
    val isLeaf : tree_t -> bool
    val nChildren : tree_t -> int
    val children : tree_t -> tree_t list
    val nthChild : (tree_t * int) -> tree_t
    val replaceNth : (tree_t * int * tree_t) -> tree_t
  end (* TREE = *)
```

The generated structure has a richer signature that is used by other grammar-dependent parts of the system.

To specify a node in a tree we use a path from the node to the tree's root. Paths and their operations are defined in a grammar-independent functor, called PathFUN, which is parameterized by a structure T matching the TREE signature. The path type is:
The path $\text{Path}(i, t, p)$ specifies node $t$, where $t$ is its parent's $i$'th child and $p$ is the path to $t$'s parent. One of the advantages of this scheme is that it promotes sharing of trees (i.e., dags), since a single physical node can have several different paths. The functor Path FUN provides functions for examining the children of a tree node that is specified by a path, tree navigation, and subtree replacement.

### 4.2 Attribution

We have chosen to store attributes and other mutable state information in auxiliary data structures. These data structures map tree nodes, specified by paths, to state information. There are a number of possible representations of these auxiliary maps; we are currently using hash tables keyed by hashing a node's path. An alternative representation that we are considering is a shadow tree.

By storing attribute instances in auxiliary structures, we gain several advantages. First, it means that the existence of an attribute instance for a particular node is completely independent of the node's representation. This allows us to trade time for space, by recomputing attribute instances instead of caching them in the tree. It also means that we can go back and forth between attributed and non-attributed trees without rewriting (although attributing a tree does require attribute evaluation).

There are two possible interfaces to the auxiliary data structures that hold the attribute instances of a tree. In the generic (or grammar-independent) approach, there is a single type of all attribute values and two functions to get and set an attribute value of a given tree node. The principal advantage of this approach is that it allows the implementation of the auxiliary state to be grammar-independent. In this approach, the code for the attribution rules must inject and project values from the attribute type. The grammar-dependent approach is to generate specialized functions for getting and setting each nonterminal's attributes. This approach is more efficient, since it exploits compile-time information to avoid run-time type discrimination. In our current prototype evaluators, which are hand-coded, we are using the former approach.

### 5 Status and Future Work

We have defined the core of our specification language and sketched the AML module facility. We have been prototyping various parts of the run-time support system (such as the tree machine and paths). We have also implemented some hand-written attribute evaluators for small grammars. We are now implementing a compiler for our specification language.

There are a number of issues that we are planning to examine in the future:

- **Address issues of input and output:** pretty printing of trees and editing. We plan to build this on top of eXene [GR91].
- **Support a modular form of attribute grammar specifications as stated previously.**
- **Our use of ML-style pattern matching in the attribution rules gives us something very close to attribute patterns [Far92]; we would like to explore this similarity.**
- **Allow the use of type constructors, such as list, in the declaration of prodtypes. One can imagine generalizing this to arbitrary parameterized prodtypes.**
- **Investigations of incremental attribute evaluation schemes with a focus on space/time tradeoffs and parallel evaluation [KG92].**
- **The implementation of real applications using AML; in particular, a programming environment for SML.**
A forthcoming technical report [EMR] provides more details and discusses many of the issues raised in this abstract.

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References


ML and Parsing—a Position Paper

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Abstract

I would like to bring together several issues connecting ML and parsing. First, the parsing of Standard ML is not simple, as the grammar is not LR(k), so the parser in an ML system must be more sophisticated than for some other modern languages. Second, ML presents an opportunity to escape from the traditional type-unsafe parser generation technique of source-code manipulation. Third, there is considerable work being done on extensible syntax in other languages (and some on quotation and anti-quotation in ML), and there is a natural and clean technique, using parsing technology, allowing the user to extend the ML syntax arbitrarily.

1 Parsing ML

The ML grammar[4] is not suited to simple parsing. The definition was written before any parser was constructed for the language, and was not inspected for any of the common desirable grammar properties (such as LALR(1)). It is not LALR(1) in several places, and not LR(k) in at least one (typed sub-patterns). Infix and precedence declarations make parsing of application expressions so awkward that this is generally handled in a post-parse phase. None of the current range of ML systems accept the full language.

There should certainly be work within the ML community to decide a new standard grammar for ML, and there has been some progress towards this. However, while we have this definition, it seems wise to work as closely to it as possible in all regards, and to build a parser that accepts the full language as defined. So we need a parsing algorithm which will accept the ML grammar. There are several such techniques [2, 7], and some recent work has been done by Jan Rekers on parser generation for these parsing algorithms[5]. However, none of these techniques have yet been applied to ML. Rekers' algorithm is particularly useful as it takes standard LR(k) parsing tables (usually k =0 or 1, for small tables) and is very efficient for grammars which are 'nearly' LR(1).
2 Parser Generation in ML

Traditional parser generators[3] are text manipulation programs, which read files written in some grammar description language, calculate (typically LALR(1)) parsing tables for the grammars, and cut-and-paste sections of 'source code' (unparsed strings) around a textual representation of the parsing tables in order to produce the source of a parser, output as pure text. This is an unsatisfactory approach, with several problems:

- one needs to define a language for describing grammars, and read (i.e. lex, parse, and error-check) raw text files in this language. The language has no computational power, and is simply a way of providing the parser generator with the hierarchical structure of a grammar and a set of associated strings for the actions. It is not even possible to provide expressions for these strings, to be evaluated at generation time, except in a very restricted sense (unless one chooses to add another layer of generation and write code to produce the original text file).

- one must write 'disembodied' sections of code, which are not compiled until after the parser generation. Compiler error messages which emerge at this stage must be traced by examining the generated source and extrapolating back to the original source error. Making corrections then requires the parser generation to be re-run.

- the output parsing code is bound together with the parse tables in a single object. One cannot change the parse tables and retain the parser without recompilation. So allowing extensible syntax is difficult.

- the 'make' actions for the grammar are completely different to those for the other parts of the language system. If one is using a specialised make system (for instance, one acting solely upon ML source files), the parser generation must be dealt with separately.

(Note: there are some sophisticated systems, such as those used at INRIA, which go some way to ameliorate the first two problems, by (for instance) providing good cross-referencing of compile-time errors and a more powerful grammar-description language allowing one to specify better error-recovery behaviour. But the basic problem is caused by dealing with source text without semantics, and that is not corrected in those systems).

ML provides an alternative approach—to define the signature of a grammar (with actions), and to write a parser generator as a functor, taking a grammar as argument and returning a parser. Then the full power of ML is available for building grammars and their actions, the type-safety of ML ensures that the generated parser does not cause runtime failures (the equivalent of the compiler errors in the previous model), a suitable structure allows different parse tables to
be exchanged for extensible syntax, and the grammar can be simply integrated into the ML separate compilation world.

Various work has been done on this in the past, particularly by Nick Rothwell at LFCS[6]. I have worked on this code, and hope to produce a more fully integrated version.

One reason why this approach has not been commonly employed before is that there are type difficulties, seemingly requiring the different non-terminals of the grammar to have a common type. The typical solution is to define a single sum type (using the datatype declaration) of all the node types of the abstract syntax. This introduces a layer of constructing and deconstructing (with exception-raising on failed matches) without adding any real type-safety, and the cost of these additional functions is high, since they must be called on every 'reduce' action of the parser. I have been working on a new approach, using simple pairs of functions which should be eliminated by a good optimizing compiler.

Such a functor approach can also be applied to lexer generation (and has been so applied), and I shall be integrating parser and lexer generation in this way (so that they share a 'token' type, and present a single interface to the rest of a compiler). Both parser and lexer generators are general, and can be applied to other languages than ML. The parsers generated use Rekers' algorithm, so can be applied to any context-free grammar.

3 Extending the ML syntax

The New Jersey compiler[1] is largely written in ML, and has lexers and parsers generated using traditional techniques. These can be replaced by functorized versions. Given a clear, public definition of the abstract syntax used by the parser, and public access to the parser generator, users can define their own concrete syntax for ML. By separating the parse tables (generated by the parser generator) from the parser itself, the user can be allowed to 'switch in' their own parse tables, including parts of their own language in their ML code, thereby extending the syntax. Specialized syntaxes can be defined for particular problems, and user code and data can be written in their own form in a type-safe, controlled way.

I have been working on the New Jersey parser with this goal, and although there are some difficulties (the parser is quite tightly integrated to the rest of the compiler, especially for error-handling), they are not insuperable.

Ultimately, this tool provides a technique for the concrete syntax of ML to be 'corrected' without losing the abstract syntax on which the semantics, which are commonly perceived as the main strength of the language, are based.
References


